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ASSAYS FOR NUCLEAR RECEPTOR LIGANDS USING FRET

Abstract:

Abstract of WO9918124

Provided is a method of identifying agonists and antagonists of nuclear receptors that comprises measuring agonist-dependent fluorescence resonance energy transfer (FRET) between a fluorescently labeled nuclear receptor or ligand binding domain and fluorescently labeled CREB-binding protein (CBP), p300, other nuclear co-activator, or binding portion thereof. The method is simple, rapid, and inexpensive. Nuclear receptors and nuclear receptor co-activators labeled with fluorescent reagents for use in the above-described method are also provided. Data supplied from the esp@cenet database - Worldwide

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#### (54) Title: ASSAYS FOR NUCLEAR RECEPTOR LIGANDS USING FRET

#### (57) Abstract

Provided is a method of identifying agonists and antagonists of nuclear receptors that comprises measuring agonist-dependent fluorescence resonance energy transfer (FRET) between a fluorescently labeled nuclear receptor or ligand binding domain and fluorescently labeled CREB-binding protein (CBP), p300, other nuclear co-activator, or binding portion thereof. The method is simple, rapid, and inexpensive. Nuclear receptors and nuclear receptor co-activators labeled with fluorescent reagents for use in the above-described method are also provided.

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#### TITLE OF THE INVENTION

ASSAYS FOR NUCLEAR RECEPTOR LIGANDS USING FRET

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# CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims the benefit of U.S. Provisional Application No. 60/061,385, filed 10/7/97, the contents of which are incorporated herein by reference in their entirety.

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STATEMENT REGARDING FEDERALLY-SPONSORED R&D Not applicable.

#### REFERENCE TO MICROFICHE APPENDIX

Not applicable.

#### FIELD OF THE INVENTION

This invention relates to methods of identifying novel agonists and antagonists of nuclear receptors utilizing the agonist-dependent interaction of such receptors with CREB-binding protein (CBP) or other nuclear receptor co-activators in which this interaction is detected by fluorescence resonance energy transfer.

# BACKGROUND OF THE INVENTION

Nuclear receptors are a superfamily of ligand-activated transcription factors that bind as homodimers or heterodimers to their cognate DNA elements in gene promoters. The superfamily, with more than 150 members, can be divided into subfamilies (e.g. the steroid, retinoid, thyroid hormone, and peroxisome proliferator-activated [PPAR] subfamilies). Each subfamily may consist of several members which are encoded by individual genes (e.g. PPARα, PPARγ, and PPARδ). In addition, alternative mRNA splicing can result in more than one isoform of these genes as in the case of specific PPARs (e.g. PPARγ1 and PPARγ2). The nuclear receptor superfamily is involved in a wide variety of physiological functions in mammalian cells: e.g., differentiation, proliferation, and metabolic homeostasis. Dysfunction

or altered expression of specific nuclear receptors has been found to be involved in disease pathogenesis.

The PPAR subfamily of nuclear receptors consists of three members: PPARα, PPARγ, and PPARδ. PPARα is highly expressed in liver and kidney. Activation of PPARa by peroxisome proliferators (including hypolipidimic reagents such as fibrates) or medium and long-chain fatty acids is responsible for the induction of acyl-CoA oxidase and hydratase-dehydrogenase (enzymes required for peroxisomal β-oxidation), as well as cytochrome P450 4A6 (an enzyme 10 required for fatty acid ω-hydroxylase). Thus, PPARα has an important role in the regulation of lipid metabolism and is part of the mechanism through which hypolipidimic compounds such as fibrates exert their effects. PPARy is predominantly expressed in adipose tissue. Recently, a prostaglandin J2 metabolite, 15-Deoxy-D12,14-prostaglandin J2, has 15 been identified as a potential physiological ligand of PPARy. Both 15-Deoxy-D12,14-prostaglandin J2 treatment of preadipocytes or retroviral expression of PPARy2 in fibroblasts induced adipocyte differentiation, demonstrating the role of PPARy in adipocyte differentiation and lipid storage. The demonstration that anti-diabetic and lipid-lowering 20 insulin sensitizing compounds known as thiazolidinediones are high affinity ligands for PPARy suggests a broad therapeutic role for PPARy ligands in the treatment of diabetes and disorders associated with insulin resistance (e.g. obesity and cardiovascular disease).

Nuclear receptor proteins contain a central DNA binding domain (DBD) and a COOH-terminal ligand binding domain (LBD). The DBD is composed of two highly conserved zinc fingers that target the receptor to specific promoter/enhancer DNA sequences known as hormone response elements (HREs). The LBD is about 200-300 amino acids in length and is less well conserved than the DBD. There are at least three functions for the LBD: dimerization, ligand binding, and transactivation. The transactivation function can be viewed as a molecular switch between a transcriptionally inactive and a transcriptionally active state of the receptor. Binding of a ligand which is an agonist flips the switch from the inactive state to the active state. The COOH-terminal portion of the LBD contains an activation function domain (AF2) that is required for the switch.

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The ligand-induced nuclear receptor molecular switch is mediated through interactions with members of a family of nuclear receptor co-activators (e.g., CBP/p300, SRC-1/NcoA-1, TIF2/GRIP-1/NcoA-2, and p/CIP). Upon binding of agonist to its cognate receptor 5 LBD, a conformational change in the receptor protein creates a coactivator binding surface and results in recruitment of co-activator(s) to the receptor and subsequent transcriptional activation. The binding of antagonist ligands to nuclear receptors will not induce the required conformational change and prevents recruitment of co-activator and subsequent induction of transcription. The co-activators CREB-binding 10 protein (CBP) and p300 are two closely related proteins that were originally discovered by virtue of their ability to interact with the transcription factor CREB. These two proteins share extensive amino acid sequence homology. CBP can form a bridge between nuclear receptors and the basic transcriptional machinery (Kamei et al., 1996, 15 Cell 85:403-414; Chakravarti et al., 1996, Nature 383:99-103; Hanstein et al., 1996, Proc. Natl. Acad. Sci. USA 93:11540-11545; Heery et al., 1997, Nature 387:733-736). CBP also contains intrinsic histone acetyltransferase activity which could result in local chromatin rearrangement and further activation of transcription. Ligand- and 20 AF2-dependent interaction between certain nuclear receptors and CBP has been demonstrated in in vitro pull down assays and far-western assays. This interaction is both necessary and sufficient for the transcriptional activation that is mediated by these nuclear receptors. Thus, an AF2 mutant of the estrogen receptor (ER) which abolishes the 25 transcriptonal function of the receptor is incapable of interacting with CBP.

The N-termini of CBP and p300 have been shown to interact with the ligand-binding domains of some nuclear receptors (Kamei et al., 1996, Cell 85:403-414, hereinafter "Kamei"). Kamei was able to demonstrate direct interaction of CBP and p300 with nuclear receptors by several different methods:

(1) Kamei produced GST fusion proteins of the first 100 amino acids of the N-terminus of CBP. These fusion proteins were run out on a polyacrylamide gel, transferred to a membrane, and the membrane was exposed to 32P-labeled ligand-binding domains of

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nuclear receptors. In the presence of ligand, a specific binding interaction between the CBP and nuclear receptor fragments was detected in that the <sup>32</sup>P-labeled ligand-binding domains were observed to bind to the bands on the membrane containing the GST-CBP fusion proteins.

- (2) Kamei also utilized the yeast two-hybrid system. The ligand-binding domain of the nuclear receptor fused to the DNA-binding domain of the LexA protein was used as bait. The amino terminal domain of CBP fused to the gal4 transactivation domain was used as prey. In the presence of ligand, a specific binding interaction (occurring in vivo, i.e., within the yeast) was observed between the CBP and nuclear receptor fragments.
- (3) Kamei observed ligand-induced binding between CBP and nuclear receptors via a gel-shift assay. This assay is based on the observation that, in the presence of ligand, nuclear receptors will bind to oligonucleotides containing their target recognition sequence. Such binding results in the formation of a nuclear receptor-ligand-oligonucleotide complex having a higher molecular weight than the oligonucleotide alone. This difference in molecular weight is detected via a shift in position of the 32P-labeled oligonucleotide when it is run out on a polyacrylamide gel. Kamei found that a fragment of CBP (the N-terminal 100 amino acids) was capable of binding to the nuclear receptor-ligand-oligonucleotide complex and shifting the complex's position on the gel to an even higher molecular weight.
- (4) Kamei was able to co-immunoprecipitate CBP using antibodies to nuclear receptors in extracts from a variety of cells in the presence of ligand.
- (5) By the use of transcriptional activation assays, Kamei was able to demonstrate that nuclear receptors and CBP interact in a functional manner. Such transcriptional activation assays can indicate that two proteins are involved in a pathway that results in transcriptional activation but these assays do not prove that the interaction between the proteins is one of direct binding.

By the above-described methods, Kamei was able to demonstrate specific binding interactions between CBP and the retinoic acid receptor (RAR), glucocorticoid receptor (GR), thyroid hormone

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receptor (T3R), and retinoid X receptor (RXR). Kamei also demonstrated specific binding between the N-terminus of p300 and RAR. However, Kamei did not demonstrate specific binding between CBP, p300, or any other nuclear receptor co-activators and PPARs.

What is striking about the methods used by Kamei is their extremely laborious and time consuming nature. Such methods involve, among other things, the construction of fusion proteins, the preparation of <sup>32</sup>P-labeled proteins, the construction of specialized expression vectors for the yeast two-hybrid assay and the transcriptional activation assays, the running of many gels, and the raising of antibodies. Most of these assays take days to carry out and preparing the reagents needed to carry them out may take weeks. Because of the complicated reagents that are involved in these assays and the time needed to prepare and run the assays, these assays tend to be costly. Investigators other than Kamei who have studied the interaction between nuclear receptors and CBP have also been forced to rely on such cumbersome methods (see, e.g., Chakravarti et al., 1996, Nature 383:99-103; Hanstein et al., 1996, Proc. Natl. Acad. Sci. USA 93:11540-11545; Heery et al., 1997, Nature 387:733-736).

Kamei did not use the above-described methods to identify novel agonists or antagonists of nuclear receptors. The focus of Kamei was not on agonists or antagonists, but rather on the interaction between nuclear receptors and CBP. Although modifying the methods of Kamei to identify agonists or antagonists might be possible, such methods would suffer from serious disadvantages. This is because, as discussed above, all of the assays employed by Kamei to study the interaction of CBP and p300 with nuclear receptors are very laborious, slow, and costly. Given the therapeutic importance of steroid hormones such as estrogen, cortisol, progesterone, and other nuclear receptor agonists such as thyroid hormone and antidiabetic thiazolidinedione compounds, the need for improved high-throughput screening assays to identify potential pharmaceutical compounds affecting nuclear receptors is clear. Historically, therapeutically useful nuclear receptor ligand compounds were identified by screening animal models, an approach which is even more labor intensive and time consuming than the methods used by Kamei. Also, approaches such as those used by

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Kamei are ill-suited for the identification of antagonists of nuclear receptors. It is now widely appreciated that antagonists of nuclear receptors can be valuable therapeutic agents. Examples of such therapeutically useful antagonists are tamoxifene, raloxifene, and RU-486.

What is needed is a high throughput, time and laborsaving, non-radioactive, inexpensive, and very reliable assay for the identification and characterization of both agonists and antagonists of nuclear receptors. Such an assay is provided by the present invention.

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# SUMMARY OF THE INVENTION

The present invention provides novel methods of identifying agonists and antagonists of nuclear receptors. The methods take advantage of the agonist-dependent binding of nuclear receptors and CBP, p300, or other nuclear receptor co-activators. In the absence of agonist, binding between the nuclear receptor and CBP, p300, or other nuclear receptor co-activators does not occur. If agonist is present, however, such binding occurs and can be detected by fluorescence resonance energy transfer (FRET) between a fluorescently-labeled nuclear receptor and fluorescently-labeled CBP, p300, or other nuclear receptor co-activator. Antagonists can be identified by virtue of their ability to prevent or disrupt the agonist-induced interaction of nuclear receptors and CBP, p300, or other nuclear receptor co-activators. In contrast to prior art methods of identifying agonists and antagonists of nuclear receptors, the methods of the present invention, are simple, rapid, and less costly.

The present invention provides a nuclear receptor or ligand binding domain thereof labeled with a fluorescent reagent for use in the above-described methods of identifying agonists and antagonists of nuclear receptors. The present invention also provides CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, labeled with a fluorescent reagent.

## BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 illustrates a method of fluorescently labelling a protein or polypeptide with Europium cryptate (Eu3+K).

Figure 2 illustrates the format for experiments 1 and 2 of Table 1.

Figure 3 illustrates the format for experiment 3 of Table 1.

Figure 4 illustrates the format for experiment 4 of

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Figure 5 shows the results of studies using the methods of the present invention with four known PPAR $\gamma$  agonists. --o-- = AD5075; -- $\square$ -- = Pioglitazone; -- $\times$ -- = Troglitazone; -- $\lozenge$ -- = BRL49653.

Figure 6 shows a measurement of the binding constant for the interaction between hCBP and PPARy1LBD.

Figure 7A shows the amino acid sequence of human CBP (SEQ.ID.NO.:1).

Figure 7B shows the nucleotide sequence of a cDNA encoding human CBP (SEQ.ID.NO.:2). The open reading frame is at positions 76-1290.

Figure 8A shows the amino acid sequence of human PPARα (SEQ.ID.NO.:3).

Figure 8B shows the nucleotide sequence of a cDNA encoding human PPAR $\alpha$  (SEQ.ID.NO.:4). The open reading frame is at positions 217-1623.

Figure 9A shows the amino acid sequence of human PPARy1 (SEQ.ID.NO.:5).

Figure 9B shows the nucleotide sequence of a cDNA encoding human PPARγ1 (SEQ.ID.NO.:6). The open reading frame is at positions 173-1609.

Figure 10A shows the amino acid sequence of human PPARδ (SEQ.ID.NO.:7).

Figure 10B-C shows the nucleotide sequence of a cDNA encoding human PPARδ (SEQ.ID.NO.:8). The open reading frame is at positions 338-1663.

# DETAILED DESCRIPTION OF THE INVENTION

For the purposes of this invention:

- an "agonist" is a substance that binds to nuclear receptors in such a way that a specific binding interaction between the nuclear receptor and CBP or other nuclear receptor co-activator can occur.

- an "antagonist" is a substance that is capable of preventing or disrupting the agonist-induced specific binding interaction between a nuclear receptor and CBP, p300, or another nuclear receptor coactivator.
- a "ligand" of a nuclear receptor is an agonist or an antagonist of the nuclear receptor.
- a "specific binding interaction," "specific binding," and the like, refers to binding between a nuclear receptor and CBP, p300, or other nuclear receptor co-activator which results in the occurrence of fluorescence resonance energy transfer between a fluorescent reagent bound to the nuclear receptor and a fluorescent reagent bound to CBP, p300, or other nuclear receptor co-activator.

With respect to CBP, p300, or other nuclear receptor coactivators, a "binding portion" is that portion of CBP, p300, or other nuclear receptor co-activators that is sufficient for specific binding interactions with nuclear receptors.

With respect to nuclear receptors, a "ligand binding domain" is that portion of a nuclear receptor that is sufficient to bind an agonist or antagonist of the nuclear receptor.

The present invention provides a high throughput, time and labor-saving, non-radioactive, inexpensive, and very reliable assay for the identification and characterization of both agonists and antagonists of nuclear receptors. In a general embodiment, the present invention provides methods of identifying agonists and antagonists for any nuclear receptor for which CBP, p300, or another nuclear receptor binding protein is a co-activator. Such agonists and antagonists are identified by virtue of their ability to induce or prevent binding between the ligand binding domain of a nuclear receptor and CBP, p300, or other nuclear receptor co-activator. The interaction between the nuclear receptor and CBP, p300, or other nuclear receptor co-activator is monitored by observing the occurrence of fluorescence resonance energy

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transfer (FRET) between two fluorescent reagents. One fluorescent reagent is bound to the nuclear receptor; the other fluorescent reagent is bound to CBP, p300, or other nuclear receptor co-activator. The binding of fluorescent reagent to nuclear receptor, CBP, p300, or other nuclear receptor co-activator can be by a covalent linkage or a non-covalent linkage.

The present invention makes use of fluorescence resonance energy transfer (FRET). FRET is a process in which energy is transferred from an excited donor fluorescent reagent to an acceptor fluorescent reagent by means of intermolecular long-range dipole-dipole coupling. FRET typically occurs over distances of about 10è to 100è and requires that the emission spectrum of the donor reagent and the absorbance spectrum of the acceptor reagent overlap adequately and that the quantum yield of the donor and the absorption coefficient of the acceptor be sufficiently high. In addition, the transition dipoles of the donor and acceptor fluorescent reagents must be properly oriented relative to one another. For a review of FRET and its applications to biological systems, see Clegg, 1995, Current Opinions in Biotechnology 6:103-110.

The present invention makes use of a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent and CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, labeled with a second fluorescent reagent. The second fluorescent reagent comprises a fluorophore capable of undergoing energy transfer by either (a) donating excited state energy to the first fluorescent reagent, or (b) accepting excited state energy from the first fluorescent reagent. In other words, according to the present invention, either the first or the second fluorescent reagents can be the donor or the acceptor during FRET.

The first and second fluorescent reagents are spectropscopically complementary to each other. This means that their spectral characteristics are such that excited state energy transfer can occur between them. FRET is highly sensitive to the distance between the first and second fluorescent reagents. For example, FRET varies inversely with the sixth power of the distance between the first and second fluorescent reagents. In the absence of agonist, the first

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fluorescent reagent, bound to the nuclear receptor or ligand binding domain thereof, will not be near the second fluorescent reagent, bound to CBP, p300, or other nuclear receptor co-activator, or binding portion thereof. Thus, no FRET, or very little FRET, will be observed. In the presence of agonist, however, interaction between the nuclear receptor and CBP, p300, or other nuclear receptor co-activator will occur, thus bringing close together the first and the second fluorescent reagents, allowing FRET to occur and be observed.

Accordingly, the present invention provides a method of identifying an agonist of a nuclear receptor that comprises providing:

- (a) a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent;
- (b) CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, labeled with a second fluorescent reagent; and
- (c) a substance suspected of being an agonist of the nuclear receptor;

under conditions such that, if the substance is an agonist of the nuclear receptor, binding between the nuclear receptor or ligand binding domain thereof and CBP, p300, or other nuclear receptor coactivator, or a binding portion thereof, will occur; and

(d) measuring fluorescence resonance energy transfer (FRET) between the first and second fluorescent reagents;

where the occurrence of FRET indicates that the substance is an agonist of the nuclear receptor.

In particular embodiments, the nuclear receptor is selected from the group consisting of steroid receptors, thyroid hormone receptors, retinoic acid receptors, peroxisome proliferator-activated receptors, retinoid X receptors, glucocorticoid receptors, vitamin D receptors, and "orphan nuclear receptors" such as LXR, FXR, etc.

In a particular embodiment, the nuclear receptor or ligand binding domain thereof is a full-length nuclear receptor. In another embodiment, the nuclear receptor or ligand binding domain thereof is a ligand binding domain of a nuclear receptor. In another embodiment, the nuclear receptor or ligand binding domain thereof comprises an AF-2 site of a nuclear receptor.

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In a particular embodiment, the nuclear receptor or ligand binding domain thereof is a full-length PPAR. In another embodiment, the nuclear receptor or ligand binding domain thereof is the ligand binding domain of a PPAR. In a further embodiment, the PPAR is selected from the group consisting of PPARα, PPARγ1, PPARγ2, and PPARδ. In a further embodiment, the ligand binding domain of the PPAR contains amino acid residues 176-478 of human PPARγ1.

In a particular embodiment, the nuclear receptor or ligand binding domain thereof contains amino acids 143-462 of human RARα.

In another embodiment, the nuclear receptor or ligand binding domain thereof contains amino acids 122-410 of rat T<sub>3</sub>Rα1. In another embodiment, the nuclear receptor or ligand binding domain thereof contains amino acids 227-463 of mouse RXRγ. In another embodiment, the nuclear receptor or ligand binding domain thereof contains amino acids 251-595 of human ER.

In a particluar embodiment, the above-described methods utilize full-length CBP, either mouse or human. In other embodiments, the methods utilize amino acid residues 1-113 of human CBP. In another embodiment, the methods utilize amino acid residues 1-453 of human CBP.

The conditions under which the methods described above are carried out are conditions that are typically used in the art for the study of protein-protein interactions: e.g., physiological pH; salt conditions such as those represented by such commonly used buffers as PBS; a temperature of about 4°C to about 55°C. The presence of commonly used non-ionic detergents, e.g., NP-40®, sarcosyl, Triton X-100®, is optional. When europium cryptates are used as fluorescent reagents, reactions should contain KF at a concentration of at least 200 mM.

Heery et al., 1997, Nature 387:733-736 showed that interactions between nuclear receptors and a variety of nuclear receptor co-activators are mediated by a short amino acid sequence in the nuclear receptor co-activators having the amino acid sequence LXXLL, where L is leucine and X represents any amino acid. Accordingly, the present invention can be practiced with a binding portion of a nuclear receptor co-activator, provided that the binding portion contains the amino acid

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sequence LXXLL. Therefore, the present invention includes a method of identifying an agonist of a nuclear receptor that comprises providing:

- (a) a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent;
- (b) a binding portion of a nuclear receptor co-activator, where the binding portion contains the amino acid sequence LXXLL, and where the binding portion is labeled with a second fluorescent reagent; and
- (c) a substance suspected of being an agonist of the nuclear receptor;

under conditions such that, if the substance is an agonist of the nuclear receptor, binding between the nuclear receptor or ligand binding domain thereof and the binding portion of the nuclear receptor co-activator will take place; and

(d) measuring fluorescence resonance energy transfer (FRET) between the first and second fluorescent reagents;

where the occurrence of FRET indicates that the substance is an agonist of the nuclear receptor.

In a particular embodiment, the nuclear receptor coactivator is selected from the group consisting of: human RIP-140, human SRC-1, mouse TIF-2, human or mouse CBP, human or mouse p300, mouse TIF-1, and human TRIP proteins.

In a particular embodiment, the nuclear receptor coactivator is human RIP-140 and the binding portion includes a contiguous stretch of amino acids of human RIP-140 selected from the group consisting of: positions 20-29, 132-139, 184-192, 266-273, 379-387, 496-506, 712-719, 818-825, 935-944, and 935-942.

In another embodiment, the nuclear receptor co-activator is human SRC-1 and the binding portion includes a contiguous stretch of amino acids of human SRC-1 selected from the group consisting of: positions 45-53, 632-640, 689-696, 748-755, and 1434-1441.

In another embodiment, the nuclear receptor co-activator is mouse TIF-2 and the binding portion includes a contiguous stretch of amino acids of mouse TIF-2 selected from the group consisting of: positions 640-650, 689-699, and 744-754.

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In another embodiment, the nuclear receptor co-activator is human or mouse CBP and the binding portion includes a contiguous stretch of amino acids of human or mouse CBP selected from the group consisting of: positions 68-78 and 356-366.

In another embodiment, the nuclear receptor co-activator is human or mouse p300 and the binding portion includes a contiguous stretch of amino acids of human or mouse p300 selected from the group consisting of: positions 80-90 and 341-351.

In another embodiment, the nuclear receptor co-activator is mouse TIF-1 and the binding portion includes a contiguous stretch of amino acids of mouse TIF-1 containing positions 722-732.

In another embodiment, the nuclear receptor co-activator is human TRIP2 and the binding portion includes a contiguous stretch of amino acids of human TRIP2 containing positions 23-33.

In another embodiment, the nuclear receptor co-activator is human TRIP3 and the binding portion includes a contiguous stretch of amino acids of human TRIP3 containing positions 97-107.

In another embodiment, the nuclear receptor co-activator is human TRIP4 and the binding portion includes a contiguous stretch of amino acids of human TRIP4 containing positions 36-46.

In another embodiment, the nuclear receptor co-activator is human TRIP5 and the binding portion includes a contiguous stretch of amino acids of human TRIP5 containing positions 26-36.

In another embodiment, the nuclear receptor co-activator is human TRIP8 and the binding portion includes a contiguous stretch of amino acids of human TRIP8 containing positions 36-46.

In another embodiment, the nuclear receptor co-activator is human TRIP9 and the binding portion includes a contiguous stretch of amino acids of human TRIP9 selected from the group consisting of: positions 73-83, 256-266 and 288-298.

For amino acid sequences of nuclear receptor co-activators, see Yao et al., 1996, Proc. Natl. Acad. Sci. USA 93:10626-10631 (SRC-1); O§ate et al., 1995, Science 270:1354-1357 (SRC-1); Cavaillès et al., 1995, EMBO J. 14:3741-3751 (RIP-140); Voegel et al., 1996, EMBO J. 15:101-108 (TIF-2); Kwok et al., 1994, Nature 370:223-226 (CBP); Arias et al., 1994, Nature 370:226-229 (CBP); Eckner et al., 1994, Genes Dev. 8:869-884

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(p300); Le Douarin et al., 1995, EMBO J. 14:2020-2033 (TIF-1); Lee et al., 1995, Nature 374:91-94 (TRIP proteins).

The particular embodiments of the present invention described above are all particular embodiments of a more general method that is also part of the present invention. That general method is a method of identifying an agonist of a nuclear receptor that comprises providing:

- (a) a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent;
- (b) a polypeptide containing the amino acid sequence LXXLL where the polypeptide is labeled with a second fluorescent reagent; and
  - (c) a substance suspected of being an agonist of the nuclear receptor;
- under conditions such that, if the substance is an agonist of the nuclear receptor, binding between the nuclear receptor or ligand binding domain thereof and the polypeptide will take place; and
  - (d) measuring fluorescence resonance energy transfer (FRET) between the first and second fluorescent reagents;
  - where the occurrence of FRET indicates that the substance is an agonist of the nuclear receptor.

In a particular embodiment, the amino acid sequence LXXLL is present in an  $\alpha$  helical portion of the polypeptide. In another embodiment, the amino acid sequence LXXLL is present in an  $\alpha$  helical portion of the polypeptide and the leucines form a hydrophobic face.

The present invention provides methods for identifying antagonists of a nuclear receptor. Such methods are based on the ability of the antagonist to prevent the occurrence of agonist-induced binding between a nuclear receptor and CBP, p300, or other nuclear receptor coactivator, or to disrupt such binding after it has occurred. Thus, the present invention provides a method for identifying antagonists of nuclear receptors that comprises providing:

- (a) a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent;
- (b) CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, labeled with a second fluorescent reagent;

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(c) an agonist of the nuclear receptor; and

(d) a substance suspected of being an antagonist of the nuclear receptor;

under conditions such that, in the absence of the substance, binding between the nuclear receptor or ligand binding domain thereof and CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof will occur; and

(e) measuring fluorescence resonance energy transfer (FRET) between the first and second fluorescent reagents when the substance is present and measuring FRET between the first and second fluorescent reagents when the substance is absent;

where the a decrease in FRET when the substance is present indicates that the substance is an antagonist of the nuclear receptor.

In particular embodiments, the nuclear receptor is selected from the group consisting of steroid receptors, thyroid hormone receptors, retinoic acid receptors, peroxisome proliferator-activated receptors, retinoid X receptors, glucocorticoid receptors, vitamin D receptors, and "orphan nuclear receptors" such as LXR, FXR, etc.

In a particular embodiment, the nuclear receptor or ligand binding domain thereof is a full-length nuclear receptor. In another embodiment, the nuclear receptor or ligand binding domain thereof is a ligand binding domain of a nuclear receptor. In another embodiment, the nuclear receptor or ligand binding domain thereof is an AF-2 site of a nuclear receptor.

In a particular embodiment, the nuclear receptor or ligand binding domain thereof is a full-length PPAR. In another embodiment, the nuclear receptor or ligand binding domain thereof is the ligand binding domain of a PPAR. In a further embodiment, the PPAR is selected from the group consisting of PPARa, PPARa, and PPARa. In a further embodiment, the ligand binding domain of the PPAR contains amino acid residues 176-478 of human PPARa.

In a particular embodiment, the nuclear receptor or ligand binding domain thereof contains amino acids 143-462 of human RAR $\alpha$ . In another embodiment, the nuclear receptor or ligand binding domain thereof contains amino acids 122-410 of rat  $T_3R\alpha 1$ . In another

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embodiment, the nuclear receptor or ligand binding domain thereof contains amino acids 227-463 of mouse RXRy. In another embodiment, the nuclear receptor or ligand binding domain thereof contains amino acids 251-595 of human ER.

In a particular embodiment, the above-described methods utilize full-length CBP, either mouse or human. In other embodiments, the methods utilize amino acid residues 1-113 of human CBP. In another embodiment, the methods utilize amino acid residues 1-453 of human CBP.

The conditions under which the methods described above are carried out are conditions that are typically used in the art for the study of protein-protein interactions: e.g., physiological pH; salt conditions such as those represented by such commonly used buffers as PBS; a temperature of about 4°C to about 55°C. The presence of commonly used non-ionic detergents, e.g., NP-40®, sarcosyl, Triton X-100®, is optional. When europium cryptates are used as fluorescent reagents, reactions should contain KF at a concentration of at least 200 mM.

In principle, one could measure FRET by monitoring either (a) a decrease in the emission of the donor fluorescent reagent following stimulation at the donor's absorption wavelength and/or (b) an increase in the emission of the acceptor reagent following stimulation at the donor's absorption wavelength. In practice, FRET is most effectively measured by emission ratioing. Emission ratioing monitors the change in the ratio of emission by the acceptor over emission by the donor. An increase in this ratio signifies that energy is being transferred from donor to acceptor and thus that FRET is occurring. Emission ratioing can be measured by employing a laser-scanning confocal microscope. Emission ratioing is preferably done by splitting the emitted light from a sample with a dichroic mirror and measuring two wavelength bands (corresponding to the donor and the acceptor emission wavelengths) simultaneously with two detectors. Alternatively, the emitted light can be sampled consecutively at each wavelength (by using appropriate filters) with a single detector. In any case, these and other methods of measuring FRET are well known in the art.

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Although a variety of donor and acceptor fluorescent reagents can be used in the practice of the present invention, preferred embodiments of the present invention make use of cryptates of fluorescent reagents as donor reagents. Inclusion of a substrate into the intramolecular cavity of a macropolycyclic ligand results in the formation of a cryptate. The macropolycyclic ligand shields the substrate from interaction with solvent and other solute molecules. If the substrate is a fluororescent reagent, formation of a cryptate may result in markedly different spectroscopic characteristics for the reagent as compared to the spectroscopic characteristics of the free reagent.

The present invention includes the use of europium (EuIII) or terbium (TbIII) cryptates as donor fluorescent reagents. Such EuIII or TbIII cryptates, as well as methods for their formation, are well known in the art. For example, see Alpha et al., 1987, Angew. Chem. Int. Ed. Engl. 26:266-267; Mathis, 1995, Clin. Chem. 41:1391-1397. A europium cryptate is formed by the inclusion of a europium ion into the intramolecular cavity of a macropolycyclic ligand which contains bipyridine groups as light absorbers. When europium cryptates are present in solution together with fluoride ions, a total shielding of the europium cryptate fluorescence is occurs. The molecular structure of a europium cryptate is shown below.

Europium cryptates can be conjugated to proteins by the use of well-known heterobifunctional reagents (see, e.g., International Patent Application WO 89/05813; Prat et al., 1991, Anal. Biochem. 195:283-289; Lopez et al., 1993, Clin. Chem. 39:196-201).

The present invention includes the use of XL665 as the acceptor fluorescent reagent. XL665 is a crosslinked derivative of allophycocyanin (APC). APC is a porphyrin containing protein which is derived from the light harvesting system of algae (Kronick, 1986, M. Immunol. Meth. 92:1-13). XL665 has an absorption maximum at =620 10 nm and an emission maximum at 665 nm. In some embodiments of the invention, XL665 is labeled with streptavidin in order to effect the binding of the streptavidin-labled XL665 to a biotin-labeled substance, e.g., CBP or the ligand binding domain of a nuclear receptor. Streptavidin labeling of XL655 and biotin labeling of CBP, or the ligand binding domain of a nuclear receptor, can be performed by well known methods.

In a preferred embodiment of the invention, XL665 as the acceptor fluorescent reagent is combined with Europium cryptate (Eu3+K) as the donor fluorescent reagent. Europium cryptate (Eu3+K) has a large Stokes shift, absorbing light at 337 nm and emitting at 620 nm. Thus, the emission maximum of Europium cryptate (Eu3+K) overlaps the absorption maximum of XL665. Europium cryptate (Eu3+K) has a large temporal shift; the time between absorption and emission of a photon is about 1 millisecond. This is advantageous because most background fluorescence signals in biological samples are short-lived. Thus the use of a fluorescent reagent such as europium cryptate, with a long fluorescent lifetime, permits time-resolved detection resulting in the reduction of background interference.

The spectral and temporal properties of europium cryptate 30 (Eu3+K) result in essentially no fluorescence background and thus assays using this fluorescent reagent can be carried out in a "mix and read" mode, greatly facilitating its use as a high throughput screening tool. For the embodiment using Europium cryptate (Eu3+K) and XL665, the measuring instrument irradiates the sample at 337 nm and 35 measures the fluorescence output at two wavelengths, 620 nm (B counts, europium fluorescence) and 665 nm (A counts, XL665 fluorescence).

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The extent of flurorescent resonance energy transfer is measured as the ratio between these two values. Typically this ratio is multiplied by 10,000 to give whole numbers.

Other FRET donor-acceptor pairs are suitable for the practice of the present invention. For example, the following donor-acceptor pairs can be used: dansyl/fluorescein; fluorescein/rhodamine; tryptophan/aminocoumarin.

The present invention provides a nuclear receptor or ligand binding domain thereof labeled with a fluorescent reagent for use in the above-described methods of identifying agonists and antagonists of nuclear receptors. The present invention also provides CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, labeled with a fluorescent reagent.

In a particular embodiment, the nuclear receptor or ligand binding domain thereof is selected from the group consisting of PPARα, PPARγ, PPARγ, a ligand binding domain of PPARα, PPARγ, or PPARδ, and amino acid residues 176-478 of human PPARγ1 and the fluorescent reagent is selected from the group consisting of XL665 and Europium cryptate (Eu3+K).

In a particular embodiment, CBP, p300, or other nuclear receptor co-activator is labeled with a fluorescent reagent selected from the group consisting of XL665 and Europium cryptate (Eu3+K).

The following non-limiting examples are presented to better illustrate the invention.

#### **EXAMPLE 1**

# Cloning, expression, and purification of human CBP and PPAR proteins

To test whether human CBP can interact with PPARs in an agonist-dependent manner, we cloned the human cDNA fragments encoding the NH2-terminal 1-113 amino acids (hCBP1-113) and 1-453 amino acids (hCBP1-453) of human CBP by the polymerase chain reaction (PCR). The DNA and amino acid sequences of human CBP are

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disclosed in Borrow et al., 1996, Nature Genet. 14:33-41 and in GenBank, accession no. U47741.

The primers used for hCBP1-113 were:

5'-ACTCGGATCCAAGCCATGGCTGAGAACTTGCTGGACGG-3'

5 (SEQ.ID.NO.:9) and

5'-CACAAAGCTTAGGCCATGTTAGCACTGTTCGG-3' (SEQ.ID.NO.: 10).

These primers were expected to amplify a 0.9 kb DNA fragment.

The primers for hCBP1-453 were:

5'-ACTCGGATCCAAGCCATGGCTGAGAACTTGCTGGACGG-3' (SEQ.ID.NO.:9) and 5'CTCAGTCGACTTATTGAATTCCACTAGCTGGAGATCC-3' (SEQ.ID.NO.:11).

These primers were expected to amplify a 1.5 kb DNA fragment..

The template for the PCR reaction was a human fetal brain cDNA library (Stratagene, Catalogue #IS 937227). Of course, any human cDNA library from a tissue expressing CBP could have been used. The PCR amplified 0.9 kb and 1.5 kp DNA fragments which were digested with restriction endonucleases and ligated into pBluescript II vector. DNA sequencing analysis confirmed that the amplified fragments were identical to the corresponding published nucleic acid sequences of human CBP.

Based on the publicly available sequences for human CBP cited above, other primers could be readily identified and prepared by those skilled in the art in order to amplify and clone other portions of cDNA encoding human CBP from appropriate cDNA libraries. Once such portions of human CBP are produced, they could be used in the methods of the present invention in a manner similar to that described herein for hCBP1-113 and hCBP1-453. The amino acid sequence of human CBP is shown in Figure 7A; the nucleic acid sequence of the cDNA encoding human CBP is shown in Figure 7B.

To express the polypeptides encoded by the PCR fragments, vectors encoding fusion proteins of the polypeptides and glutathione Stransferase (GST) were constructed and expressed in *E. coli*. The PCR fragments were subcloned into the expression vector pGEX (Pharmacia Biotech) to generate pGEXhCBP1-113 and pGEXhCBP1-453.

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pGEXhCBP1-113 and pGEXhCBP1-453 were transfected into the DH5\alpha strain of E. coli (GIBCO BRL) and the bacteria hosting either pGEXhCBP1-113 or pGEXhCBP1-453 were cultured in LB medium (GIBCO BRL) to a density of  $OD_{600} = 0.7-1.0$  and induced for 5 overexpression of the GST-CBP fusion proteins by addition of IPTG (isopropylthio-β-galactoside) to a final concentration of 0.2 mM. The IPTG induced cultures were further grown at room temperature for 2-5 hrs. The cells were harvested by centrifugation for 10 min at 5000g. The cell pellet was used for GST-CBP fusion protein purification by following the procedure from Pharmacia Biotech using Glutathione Sepharose 10 beads. hCBP1-113 and hCBP1-453 proteins were generated by cleaving the corresponding GST fusion proteins with thrombin. SDSpolyacrylamide gel electrophoresis analysis showed that the preparation from pGEXhCBP1-113 gave two polypeptide bands, with apparent 15 molecular weight of 12 kd and 10 kd. The 12 kd band is the expected size of hCBP1-113 and the 10 kd band is most likely a premature translational termination product. The preparation from pGEXhCBP1-450 gave a single band with the expected size, 50 kd.

cDNAs encoding full-length PPARa and PPARy1 were subcloned into pGEX vectors for the production of GST-PPARa and GST-20 PPARyl fusion proteins in E.coli. PPARyl was cloned from a human fat cell cDNA library (see Elbrecht et al., 1996, Biochem. Biophys. Res. Comm. 224:431-437). A cDNA encoding the human PPARy1 ligand binding domain (PPARy1LBD; amino acids 176-478 of PPARy1) was subcloned from a modified pSG5 vector as a Xho I (site located in the N-25 terminus of the LBD)/ Xba I (site located in the pSG5 vector) fragment. The Xba I site was blunt-ended with T4 DNA polymerase. The 1.1 kb fragment containing the LBD was purified from an agarose gel and ligated into pGEX-KG (see Guan & Dixon, 1991, Anal. Biochem. 192:262-267) that had been digested with Xho I and Hind III (the Hind III site 30 had been blunt-ended with T4 DNA polymerase). This construct was used for the production of GST-hPPARy1LBD and hPPARy1LBD (the ligand binding domain cleaved free of GST). The overexpression and purification of PPARa, PPARy1, and PPARy1LBD were as described 35 above for CBP.

The DNA and amino acid sequences of human PPAR $\alpha$  are disclosed in Schmidt et al., 1992, Mol. Endocrinol. 6:1634-1641 and in GenBank, accession no. L07592. See Figure 8A and 8B.

The DNA and amino acid sequences of human PPARγ1 are disclosed in Greene et al., 1995, Gene Expr. 4:281-299; Qi et al., 1995, Mol. Cell. Biol. 15:1817-1825; Elbrecht et al., 1996, Biochem. Biophys. Res. Comm. 224:431-437; and in GenBank, accession no. L40904. See Figure 9A and 9B. Human PPARγ2 contains the same amino acid sequence as human PPARγ1 except for an amino terminal addition of 24 amino acids (see Elbrecht et al., 1996, Biochem. Biophys. Res. Comm. 224:431-437). Thus, the amino acid sequence of the ligand binding domain of human PPARγ2 is the same as the amino acid sequence of the ligand binding domain of human PPARγ1, although the numbering of the amino acids differs (176-478 for human PPARγ1 and 200-502 for human PPARγ2).

The DNA and amino acid sequences of human PPARδ are disclosed in Sher et al., 1993, Biochemistry 32:5598-5604 and in GenBank, accession no. L02932. See Figure 10A-C.

#### **EXAMPLE 2**

# 20 Interaction between PPARs and hCBP fragments

Experiments were first conducted using hCBP1-113 and hPPARγ1LBD. Purified hPPARγ1LBD was biotinylated with Sulfo-NHS-LC-Biotin (PIERCE) to a biotin:hPPARγ1LBD ratio of 3:1 according to the procedure provided by PIERCE. Purified hCBP1-113 was directly labeled with europium cryptate (Eu3+K) by the method illustrated in Figure 1. Biotin-labeled hPPARγ1LBD, Eu3+K-labeled hCBP1-113, and streptavidin-labeled XL665 (SA-XL665; from PACKARD) were incubated together in the presence or absence of 1 μM of known PPARγ agonist (BRL49653 or AD5075).

Thus, this experimental format made use of the fluorescent reagent pair europium cryptate (Eu3+K), which acted as donor, and XL665, which acted as acceptor. hCBP1-113 was directly labeled with europium cryptate (Eu3+K); hPPAR $\gamma$ 1LBD was indirectly labeled with XL665 by means of a biotin-streptavidin link. The emission maximum

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of europium cryptate (Eu3+K) overlaps with the absorption maximum of XL665. Therefore, when europium cryptate (Eu3+K) and XL665 are in close proximity, and the sample is illuminated with light at 337 nm (the absorption maximum of europium cryptate (Eu3+K)), FRET can occur 5 between europium cryptate (Eu3+K) and XL665. This FRET manifests itself as increased emission at 665 nm by XL665. Figure 2 shows a schematic of the format used in this experiment (experiment 1 of Table 1). When agonist is bound to hPPARy1LBD, a specific interaction occurs between hPPARy1LBD and hCBP1-113, thus bringing europium cryptate 10 (Eu3+K) and XL665 into close enough proximity for FRET to occur. In the absence of agonist, no interaction occurs between hPPARy1LBD and hCBP1-113 and thus europium cryptate (Eu3+K) and XL665 are not brought into close proximity and no FRET occurs. When FRET occurs, the amount of light given off by the sample at the emission maximum of 15 XL665 (665 nm) is increased relative to the amount of light given off by the sample at the emission maximum of europium cryptate (Eu3+K) (620 nm). Therefore, measuring the ratio of emission at 665 nm to 620 nm in the presence and the absence of a substance suspected of being an agonist allows for the determination of whether that substance actually 20 is an agonist. If the substance is an agonist, an increase in the ratio of emission at 665 nm to 620 nm in the presence of the substance will be observed.

Reactions were carried out in microtiter plates. Reaction conditions were: appropriate volume (total 250 µl) of the reaction buffer (either PBS or HEPES, see below, containing 500 mM KF, 0.1% bovine serum albumin, BSA) was added to each well, followed by addition of ligands (BRL49653 or AD5075 at a final concentration of 1 µM and 0.1% dimethylsulfoxide (DMSO) or vehicle control (0.1% DMSO), Eu3+K labeled hCBP (100 nM), biotin-hPPAR71LBD (100 nM), and streptavidin-labeled XL665 (100 nM) to appropriate wells. After mixing, 200 µl of reaction mixture was transferred to a new well. The plate was either directly measured for fluorescence resonance energy transfer (FRET) or covered with sealing tape (PACKARD) to avoid evaporation and incubated at room temperature for up to 24 hrs before measuring FRET.

The results of this experiment and others described below yielded ratio values as follows:

Table 1

Experiment	Buffer	Emission ratio with AD5075	Emission ratio with vehicle
1	PBS	1134	1074
2	HEPES + 0.05% NP40	967	617
3	HEPES + 0.05% NP40	1078	536
4	HEPES + 0.05% CHAPS	1883	487

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Experiment 1 of Table 1 was carried out using PBS (137 mM NaCl, 2.7 mM KCl, 4.3 mM Na2HPO4, 1.4 mM KH2PO4, pH 7.4). The greater emission ratio observed in the presence of AD5075 demonstrated that a specific interaction between hCBP1-113 and hPPARγ1LBD occurred in the presence of the agonist AD5075. Although it was clear that FRET was occurring, the signal-noise ratio was small. In experiment 2 of Table 1, HEPES buffer (N-2-hydroxyethylpiperazine-N-2-ethane sulfonic acid, 100 mM, pH 7.0) containing 0.05% NP40 (Nonidet P-40) was used instead of PBS and an improved signal-noise ratio was obtained.

In order to get an even better signal-noise ratio, the above-described format was modified slightly for experiment 3. In experiment 3, SA-XL665 (500 nM), biotin-labeled hPPAR $\gamma$ 1LBD (100 nM), GST-hCBP1-113, and Eu3+K labeled anti-GST antibody (2.5  $\mu$ l) were incubated in the presence or absence of AD5075 (1  $\mu$ M) in HEPES buffer containing 0.05% NP40. A two-fold signal-noise ratio was obtained. Figure 3 shows a schematic of the format used in experiment 3.

The anti-GST antibody was a goat antibody to GST from Pharmacia (catalogue number 27-4577-01) that was labeled with Eu3+K according to the procedure summarized below.

- Make up @ 10 mg/mL in H2O.
Need 42.2 μg (4.2 μL, 96.6 nmol) for 49.0 μg Eu3+
Reagent

$$Eu^{3+}$$

$$NH_2$$

$$NaO_3S$$

$$V$$

$$O$$

$$FW = 436.4$$

-Resuspended @ 2.5 mg/mL in 10% DMF/PBS

-FW = 1465 Use 49.0 μg (19.6 μL, 33.4 nmol) 2.9 Equiv SULFO-SMCC, 20 mM Pi buffer, 10% DMF RT, 30 minutes

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From Pharmacia, 5.0 mg/mL, FW = 150 kD Use 200 µL (1 mg, 6.66 nmol) exchange into 10 mM Borate, 350 mM NaCl, 10% Gly, pH 8.5 with BioSpin-30

5.0 Equiv SPDP, RT, 5 hours

FW =312, Dissolve @1.00 mg/mL in EtOH. Add 10.4  $\mu$ L (5 equiv., 10.4  $\mu$ g, 33.4 nmol) to protein.

To further improve the signal to noise ratio, a series of experiments were conducted. Experiment 4 of Table 1 exemplifies

results obtained from those efforts. cDNA encoding a longer fragment of hCBP was cloned and expressed to get hCBP1-453. hCBP1-453 was biotinylated. Biotin-labeled hCBP1-453 (25 nM), SA-XL665 (100 nM), GST-hPPARylLBD (1 nM), and Eu3+K-labeled anti-GST antibody (2 nM) were mixed together in the presence or absence of 1 µM AD5075. The detergent was changed from 0.05% NP40 to 0.5% CHAPS (3-{[3-cholamidopropyl]dimethyl-ammoniol}-1-propanesulfonate). A three- to

four-fold signal-noise ratio was obtained. Figure 4 shows the strategy used for experiment 4 and similar experiments.

The correlation between results from the above-described assays and previously reported results from in vitro binding and transcriptional activation assays of selected antidiabetic insulin sensitizers that are known to be PPARγ agonists (Elbrecht et al., 1996, Biochem Biophys Res Comm 224:431-437) was analyzed by titrating those known PPARγ agonists in the assays described above and comparing EC50s so obtained with previously described values for potency in

binding or transcriptional activation assays for the known agonists. The results are shown in Figure 5. From Figure 5, the following EC50s can be derived:

AD5075 = 8 nM

BRL49653 = 53 nM

Troglitazone = 646 nM

Pioglitazone = 890 nM.

These EC50s generated in the above-described assays are in close agreement with those generated by *in vitro* binding and transcriptional activation studies (Elbrecht et al., 1996, Biochem Biophys Res Comm 224:431-437).

The above-described assay can also be used to characterize the interaction between nuclear receptors with co-activators as, e.g., by determining the binding constant for that interaction. Figure 6 shows an example of such an application. Saturating amounts of PPARy agonist (10 µM BRL49653) were used. Increasing concentrations of non-biotinylated hCBP1-453 were used to titrate away biotin-hCBP-PPARy1LBD complex and decrease the fluorescence energy transfer. A Kd of 300 nM for the interaction between hCBP1-453 and PPARy1LBD can be derived from the results illustrated in Figure 6 and this Kd (300 nM) is a measurement ofthe affinity between CBP and PPARy.

The present invention is not to be limited in scope by the specific embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description. Such modifications are intended to fall within the scope of the appended claims.

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Various publications are cited herein, the disclosures of which are incorporated by reference in their entireties.

#### WHAT IS CLAIMED:

- 1. A method of identifying an agonist of a nuclear receptor that comprises providing:
- (a) a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent;
  - (b) CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, labeled with a second fluorescent reagent; and
  - (c) a substance suspected of being an agonist of the nuclear receptor;
- under conditions such that, if the substance is an agonist of the nuclear receptor, binding between the nuclear receptor or ligand binding domain thereof and CBP, p300, or other nuclear receptor coactivator, or a binding portion thereof, will occur; and
  - (d) measuring fluorescence resonance energy transfer (FRET) between the first and second fluorescent reagents;

where the occurrence of FRET indicates that the substance is an agonist of the nuclear receptor.

- 2. The method of claim 1 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of steroid receptors, thyroid hormone receptors, retinoic acid receptors, peroxisome proliferator-activated receptors, retinoid X receptors, glucocorticoid receptors, vitamin D receptors, LXR, and FXR.
- 3. The method of claim 1 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of a full-length nuclear receptor, a ligand binding domain of a nuclear receptor, and an AF-2 site of a nuclear receptor.
- 4. The method of claim 1 where the nuclear receptor or ligand binding domain thereof comprises an AF-2 site of a nuclear receptor.
- 5. The method of claim 1 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of a

full-length PPAR, a ligand binding domain of a PPAR, and amino acid residues 176-478 of human PPARy1.

- 6. The method of claim 1 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of PPARα, PPARγ1, PPARγ2, and PPARδ.
- 7. The method of claim 1 where the nuclear receptor or ligand binding domain thereof comprises a ligand binding domain selected from the group consisting of amino acids 143-462 of human RARα, amino acids 122-410 of rat T3Rα1, amino acids 227-463 of mouse RXRγ, and amino acids 251-595 of human ER.
- 8. The method of claim 1 where CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof is selected from the group consisting of full-length human CBP, full-length mouse CBP, amino acid residues 1-113 of human CBP, and amino acid residues 1-453 of human CBP.
- 9. The method of claim 1 where the first fluorescent reagent is selected from the group consisting of XL665 and Europium cryptate (Eu3+K).
- 10. The method of claim 1 where the second fluorescent reagent is selected from the group consisting of XL665 and Europium cryptate (Eu3+K).
  - 11. A method of identifying an agonist of a nuclear receptor that comprises providing:
- (a) a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent;
  - (b) a binding portion of a nuclear receptor co-activator, where the binding portion contains the amino acid sequence LXXLL, and where the binding portion is labeled with a second fluorescent
- 35 reagent; and

(c) a substance suspected of being an agonist of the nuclear receptor;

under conditions such that, if the substance is an agonist of the nuclear receptor, binding between the nuclear receptor or ligand binding domain thereof and the binding portion of the nuclear receptor co-activator will take place; and

- (d) measuring fluorescence resonance energy transfer (FRET) between the first and second fluorescent reagents;
- where the occurrence of FRET indicates that the substance is an agonist of the nuclear receptor.
  - 12. The method of claim 11 where the binding portion of a nuclear receptor co-activator is selected from the group consisting of human RIP-140, human SRC-1, mouse TIF-2, human or mouse CBP, human or mouse p300, mouse TIF-1, and human TRIP proteins.
  - 13. A method of identifying an agonist of a nuclear receptor that comprises providing:
- (a) a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent;
  - (b) a polypeptide containing the amino acid sequence LXXLL where the polypeptide is labeled with a second fluorescent reagent; and
- (c) a substance suspected of being an agonist of the nuclear receptor;

under conditions such that, if the substance is an agonist of the nuclear receptor, binding between the nuclear receptor or ligand binding domain thereof and the polypeptide will take place; and

- (d) measuring fluorescent resonance energy transfer

  (FRET) between the first and second fluorescent reagents;

  where the occurrence of FRET indicates that the substance is an agonist of the nuclear receptor.
- 14. A method for identifying an antagonist of a nuclear receptor that comprises providing:

(a) a nuclear receptor or ligand binding domain thereof labeled with a first fluorescent reagent;

- (b) CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, labeled with a second fluorescent reagent;
  - (c) an agonist of the nuclear receptor; and
- (d) a substance suspected of being an antagonist of the nuclear receptor;

under conditions such that, in the absence of the substance, binding between the nuclear receptor or ligand binding domain thereof and CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof will occur; and

(e) measuring fluorescence resonance energy transfer (FRET) between the first and second fluorescent reagents when the substance is present and measuring FRET between the first and second fluorescent reagents when the substance is absent;

where the a decrease in FRET when the substance is present indicates that the substance is an antagonist of the nuclear receptor.

15. The method of claim 14 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of steroid receptors, thyroid hormone receptors, retinoic acid receptors, peroxisome proliferator-activated receptors, retinoid X receptors, glucocorticoid receptors, vitamin D receptors, LXR, and FXR.

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16. The method of claim 14 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of a full-length nuclear receptor, a ligand binding domain of a nuclear receptor, and an AF-2 site of a nuclear receptor.

- 17. The method of claim 14 where the nuclear receptor or ligand binding domain thereof comprises an AF-2 site of a nuclear receptor.
- 18. The method of claim 14 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of a

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full-length PPAR, a ligand binding domain of a PPAR, and amino acid residues 176-478 of human PPARy1.

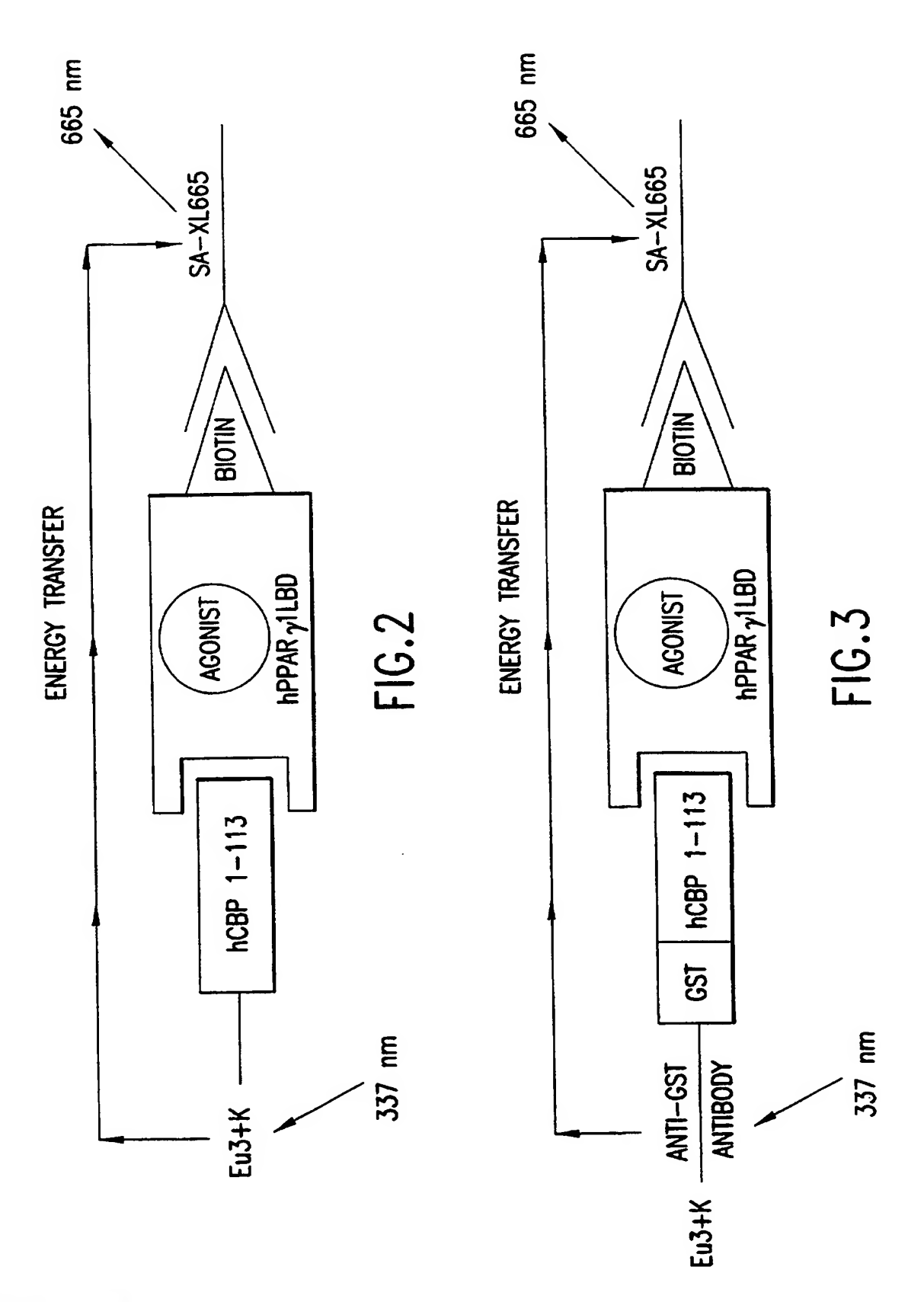
- 19. The method of claim 14 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of PPARα, PPARγ1, PPARγ2, and PPARδ.
- 20. The method of claim 14 where the nuclear receptor or ligand binding domain thereof comprises a ligand binding domain selected from the group consisting of amino acids 143-462 of human RARα, amino acids 122-410 of rat T<sub>3</sub>Rα1, amino acids 227-463 of mouse RXRγ, and amino acids 251-595 of human ER.
- 21. The method of claim 14 where CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof is selected from the group consisting of full-length CBP, amino acid residues 1-113 of human CBP, and amino acid residues 1-453 of human CBP.
- 22. The method of claim 14 where the first fluorescent reagent is selected from the group consisting of XL665 and Europium cryptate (Eu3+K).
- 23. The method of claim 14 where the second fluorescent reagent is selected from the group consisting of XL665 and Europium cryptate (Eu3+K).
  - 24. A nuclear receptor or ligand binding domain thereof labeled with a fluorescent reagent.
- 25. The nuclear receptor or ligand binding domain thereof of claim 24 where the nuclear receptor or ligand binding domain thereof is selected from the group consisting of PPARα, PPARγ1, PPARγ2, PPARδ, a ligand binding domain of PPARα, PPARγ1, PPARγ2, or PPARδ, and amino acid residues 176-478 of human PPARγ1 and the fluorescent reagent is selected from the group consisting of XL665 and Europium cryptate (Eu3+K).

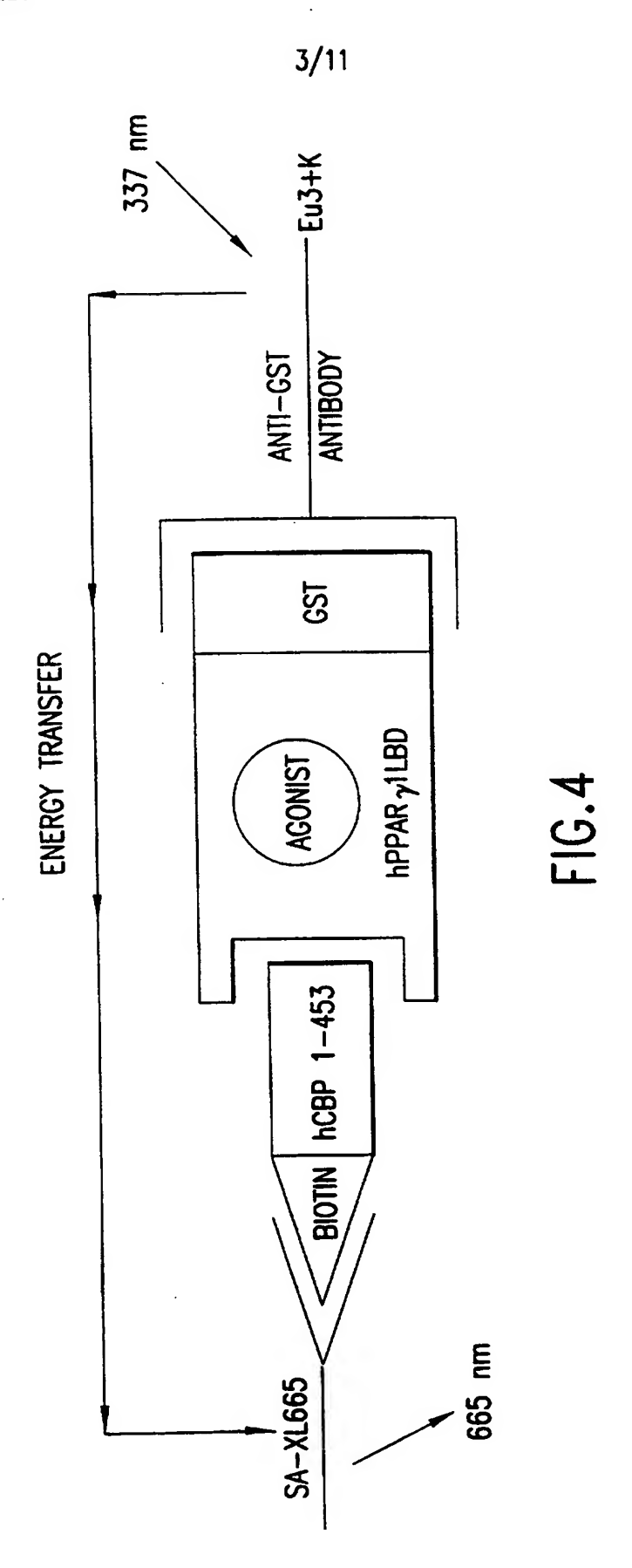
26. CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, labeled with a fluorescent reagent.

The CBP, p300, or other nuclear receptor co-activator, or a binding portion thereof, of claim 26 where the fluorescent reagent is selected from the group consisting of XL665 and Europium cryptate (Eu3+K).

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FIG.1





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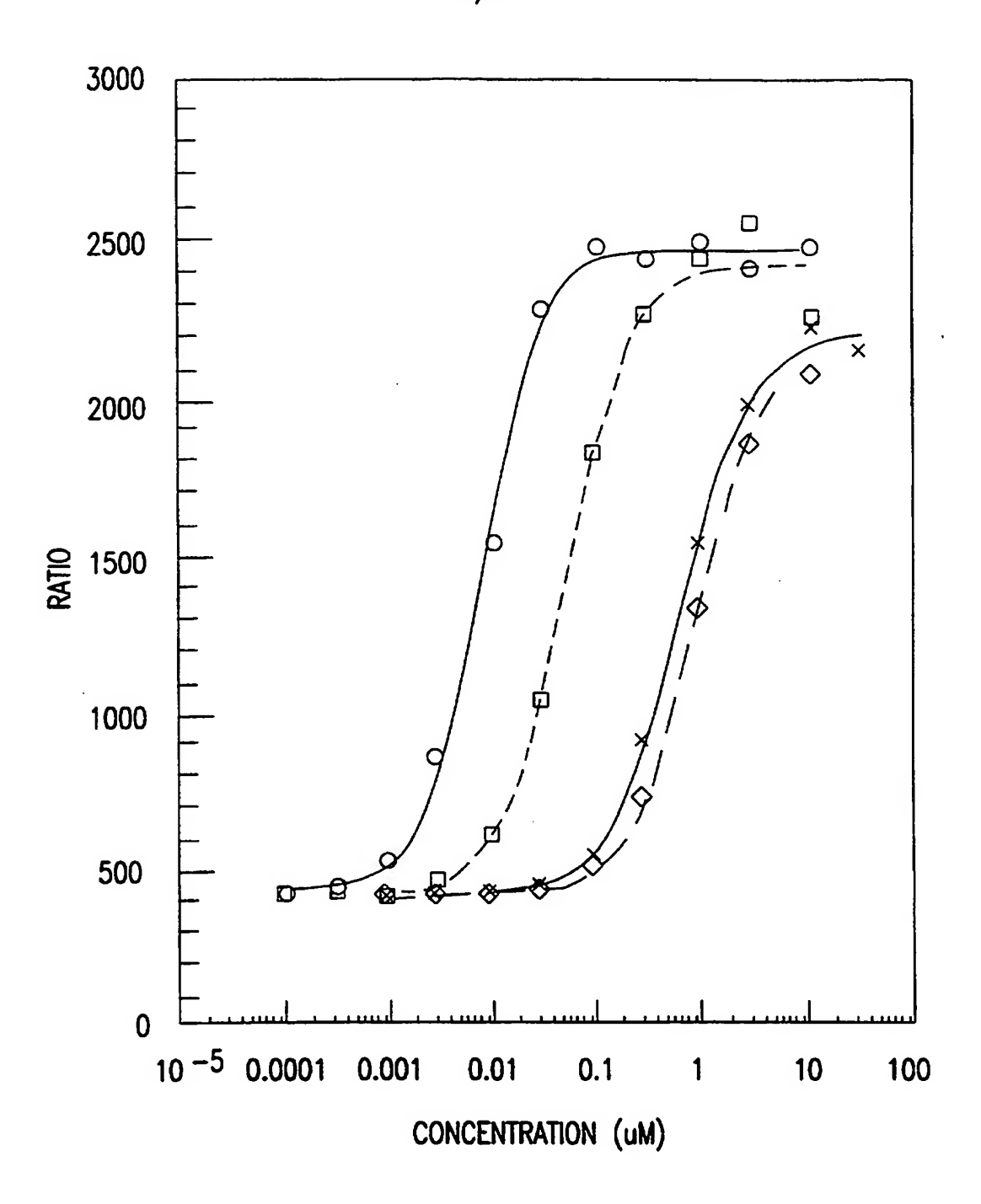


FIG.5

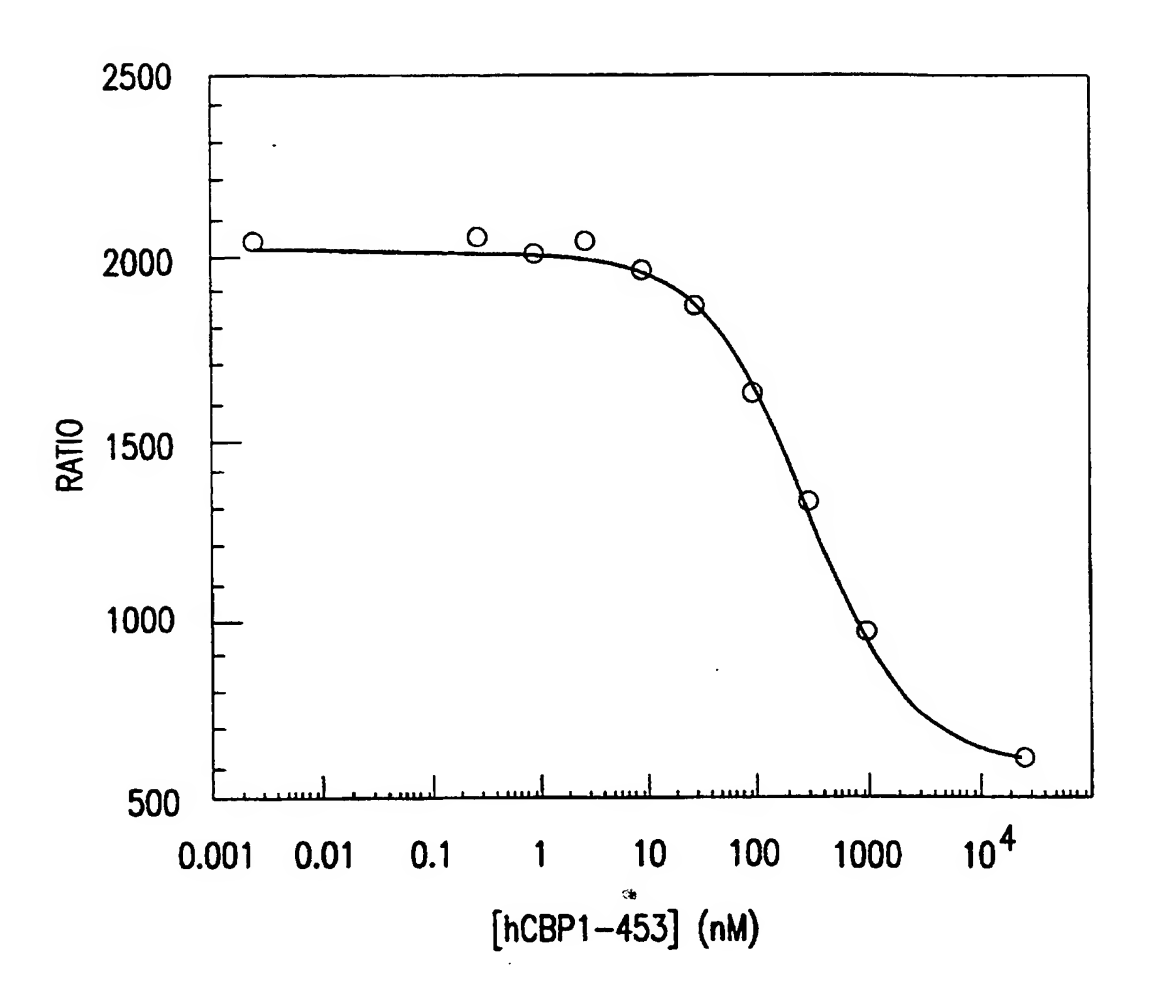


FIG.6

WO 99/18124 PCT/US98/21049 .

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- 1 MAENLLDGPPNPKRAKLSSPGFSANDSTDFGSLFDLENDLPDELIPNGGELGLLNSGNLV
- 61 PDAASKHKQLSELLRGGSGSSINPGIGNVSASSPVQQGLGGQAQGQPNSANMASLSAMGK
- 121 SPLSQGDSSAPSLPKQAASTSGPTPAASQALNPQAQKQVGLATSSPATSQTGPGICMNAN
- 181 FNQTHPGLLNSNSGHSLINQASQGQAQVMNGSLGAAGRGRGAGMPYPTPAMQGASSSVLA
- 241 ETLTQVSPQMTGHAGLNTAQAGGMAKMGITGNTSPFGQPFSQAGGQPMGATGVNPQLASK
- 301 QSMVNSLPTFPTDIKNTSVTNVPNMSQMQTSVGIVPTQAIATGPTADPEKRKLIQQQLVL
- 361 LLHAHKCQRREQANGEVRACSLPHCRTMKNVLNHMTHCQAGKACQ

# FIG.7A

```
cgagccccga ccccgtccg ggccctcgcc ggccgcgccg cccgtgcccg gggctgtttt
     cccgagcagg tgaaaatggc tgagaacttg ctggacggac cgcccaaccc caaaagagcc
61
     aaactcagct cgcccggttt ctcggcgaat gacagcacag attttggatc attgtttgac
121
    ttggaaaatg atcttcctga tgagctgata cccaatggag gagaattagg ccttttaaac
181
     agtgggaacc ttgttccaga tgctgcttcc aaacataaac aactgtcgga gcttctacga
241
     ggaggcagcg gctctagtat caacccagga ataggaaatg tgagcgccag cagccccgtg
301
     cagcagggcc tgggtggcca ggctcaaggg cagccgaaca gtgctaacat ggccagcctc
361
     agtgccatgg gcaagagccc tctgagccag ggagattctt cagcccccag cctgcctaaa
421
     caggcagcca gcacctctgg gcccaccccc gctgcctccc aagcactgaa tccgcaagca
481
     caaaagcaag tggggctggc gactagcagc cctgccacgt cacagactgg acctggtatc
541
     tgcatgaatg ctaactttaa ccagacccac ccaggcctcc tcaatagtaa ctctggccat
601
   agcttaatta atcaggcttc acaagggcag gcgcaagtca tgaatggatc tcttggggct
     gctggcagag gaaggggagc tggaatgccg taccctactc cagccatgca gggcgcctcg
     agcagcgtgc tggctgagac cctaacgcag gtttccccgc aaatgactgg tcacgcggga
781
     ctgaacaccg cacaggcagg aggcatggcc aagatgggaa taactgggaa cacaagtcca
841
     tttggacagc cctttagtca agctggaggg cagccaatgg gagccactgg agtgaacccc
901
     cagttagcca gcaaacagag catggtcaac agtttgccca ccttccctac agatatcaag
961
1021 aatacttcag tcaccaacgt gccaaatatg tctcagatgc aaacatcagt gggaattgta
1081 cccacacaag caattgcaac aggccccact gcagatcctg aaaaacgcaa actgatacag
1141 cagcagetgg ttctactgct tcatgctcat aagtgtcaga gacgagagca agcaaacgga
1201 gaggttcggg cctgctcgct cccgcattgt cgaaccatga aaaacgtttt gaatcacatg
1261 acgcattgtc aggctgggaa agcctgccaa
```

FIG.7B

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MVDTESPLCPLSPLEAGDLESPLSEEFLQEMGNIQEISQSIGEDSSGSFGFTEYQYLGSC PGSDGSVITDTLSPASSPSSVTYPVVPGSVDESPSGALNIECRICGDKASGYHYGVHACE 121 GCKGFFRRTIRLKLVYDKCDRSCKIQKKNRNKCQYCRFHKCLSVGMSHNAIRFGRMPRSE 181 KAKLKAEILTCEHDIEDSETADLKSLAKRIYEAYLKNFNMNKVKARVILSGKASNNPPFV 241 IHDMETLCMAEKTLVAKLVANGIQNKEVEVRIFHCCQCTSVETVTELTEFAKAIPAFANL 301 DLNDQVTLLKYGVYEAIFAMLSSVMNKDGMLVAYGNGFITREFLKSURKPFCDIMEPKFD 361 FAMKFNALELDDSDISLFVAAIICCGDRPGLLNVGHIEKMQEGIVHVURLHLQSNHPDDI

421 FLPKLLOKMADLRQLVTEHAQLVQIIKKTESDAALHPLLQEIYRDMY

# FIG.8A

ggcccaggct gaagctcagg gccctgtctg ctctgtggac tcaacagttt gtggcaagac aagctcagaa ctgagaagct gtcaccacag ttctggaggc tgggaagttc aagatcaaag 61 tgccagcaga ttcagtgtca tgtgaggacg tgcttcctgc ttcatagata agagtagctt 121 ggagctcggc ggcacaacca gcaccatctg gtcgcgatgg tggacacgga aagcccactc 181 tgcccctct ccccactcga ggccggcgat ctagagagcc cgttatctga agagttcctg 241 caagaaatgg gaaacatcca agagatttcg caatccatcg gcgaggatag ttctggaagc 301 tttggcttta cggaatacca gtatttagga agctgtcctg gctcagatgg ctcggtcatc 361 acggacacgc tttcaccagc ttcgagcccc tcctcggtga cttatcctgt ggtccccggc 421 agcgtggacg agtctcccag tggagcattg aacatcgaat gtagaatctg cggggacaag 481 gcctcaggct atcattacgg agtccacgcg tgtgaaggct gcaagggctt ctttcggcga 541 acgattcgac tcaagctggt gtatgacaag tgcgaccgca gctgcaagat ccagaaaaag 601 aacagaaaca aatgccagta ttgtcgattt cacaagtgcc tttctgtcgg gatgtcacac 661 aacgcgattc gttttggacg aatgccaaga tctgagaaag caaaactgaa agcagaaatt 721 cttacctgtg aacatgacat agaagattct gaaactgcag atctcaaatc tctggccaag 781 agaatctacg aggcctactt gaagaacttc aacatgaaca aggtcaaagc ccgggtcatc 841 ctctcaggaa aggccagtaa caatccacct tttgtcatac atgatatgga gacactgtgt 901 atggctgaga agacgctggt ggccaagctg gtggccaatg gcatccagaa caaggaggtg 961 1021 gaggtccgca tctttcactg ctgccagtgc acgtcagtgg agaccgtcac ggagctcacg 1081 gaattcgcca aggccatccc agcgttcgca aacttggacc tgaacgatca agtgacattg 1141 ctaaaatacg gagtttatga ggccatattc gccatgctgt cttctgtgat gaacaaagac 1201 gggatgctgg tagcgtatgg aaatgggttt ataactcgtg aattcctaaa aagcctaagg 1261 aaaccgttct gtgatatcat ggaacccaag tttgattttg ccatgaagtt caatgcactg 1321 gaactggatg acagtgatat ctcccttttt gtggctgcta tcatttgctg tggagatcgt 1381 cctgqccttc taaacgtagg acacattgaa aaaatgcagg agggtattgt acatgtgctc 1441 agactccacc tgcagagcaa ccacccggac gatatctttc tcttcccaaa acttcttcaa 1501 aaaatggcag acctccggca gctggtgacg gagcatgcgc agctggtgca gatcatcaag 1561 aagacggagt cggatgctgc gctgcacccg ctactgcagg agatctacag ggacatgtac 1621 tgagttcctt cagatcagcc acaccttttc caggagttct gaagctgaca gcactacaaa 1681 ggagacgggg gagcagcacg attttgcaca aatatccacc actttaacct tagagcttgg 1741 acagtctgag ctgtaggtaa ccggcatatt attccatatc tttgttttaa ccagtacttc 1801 taagagcata gaactcaaat gctgggggag gtggctaatc tcaggactgg gaag

FIG.8B

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- 1 MTMVDTEIAFWPTNFGISSVDLSVMEDHSHSFDIKPFTTVDFSSISTPHYEDIPFTRTDP 61 VVADYKYDLKLQEYQSAIKVEPASPPYYSEKTQLYNKPHEEPSNSLMAIECRVCGDKASG
- 121 FHYGVHACEGCKGFFRRTIRLKLIYDRCDLNCRIHKKSRNKCQYCRFQKCLAVGMSHNAI
- 181 RFGRIAQAEKEKLLAEISSDIDQLNPESADLRQALAKHLYDSYIKSFPLTKAKARAILTG
- 241 KTTDKSPFVIYDMNSLMMGEDKIKFKHITPLQEQSKEVAIRIFQGCQFRSVEAVQEITEY
- 301 AKSIPGFVNLDLNDQVTLLKYGVHEIIYTMLASLMNKDGVLISEGQGFMTREFLKSLRKP
- 361 FGDFMEPKFEFAVKFNALELDDSDLAIFIAVIILSGDRPGLLNVKPIEDIQDNLLQALEL
- 421 QLKLNHPESSQUAKLLQKMTDLRQIVTEHVQLLQVIKKTETDMSLHPLLQEIYKDLY

# FIG.9A

ccgaccttac cccaggcggc cttgacgttg gtcttgtcgg caggagacag caccatggtg ggttctctct gagtctggga attcccgagc ccgagccgca gccgccgcct ggggggcttg 61 ggtcggcctc gaggacaccg gagaggggcg ccacgccgcc gtggccgcag aaatgaccat 121 ggttgacaca gagatcgcat tctggcccac caactttggg atcagctccg tggatctctc 181 cgtaatggaa gaccactccc actcctttga tatcaagccc ttcactactg ttgacttctc 241 cagcatttct actccacatt acgaagacat tccattcaca agaacagatc cagtggttgc 301 agattacaag tatgacctga aacttcaaga gtaccaaagt gcaatcaaag tggagcctgc 361 atctccacct tattattctg agaagactca gctctacaat aagcctcatg aagagccttc 421 caactccctc atggcaattg aatgtcgtgt ctgtggagat aaagcttctg gatttcacta 481 tggagttcat gcttgtgaag gatgcaaggg tttcttccgg agaacaatca gattgaagct 541 tatctatgac agatgtgatc ttaactgtcg gatccacaaa aaaagtagaa ataaatgtca 601 gtactgtcgg tttcagaaat gccttgcagt ggggatgtct cataatgcca tcaggtttgg 661 gcggatcgca caggccgaga aggagaagct gttggcggag atctccagtg atatcgacca 721 gctgaatcca gagtccgctg acctccgtca ggccctggca aaacatttgt atgactcata 781 cataaagtcc ttcccgctga ccaaagcaaa ggcgagggcg atcttgacag gaaagacaac 841 agacaaatca ccattcgtta tctatgacat gaattcctta atgatgggag aagataaaat 901 caagttcaaa cacatcaccc ccctgcagga gcagagcaaa gaggtggcca tccgcatctt 961 tcagggctgc cagtttcgct ccgtggaggc tgtgcaggag atcacagagt atgccaaaag 1021 cattcctggt tttgtaaatc ttgacttgaa cgaccaagta actctcctca aatatggagt 1081 ccacgagatc atttacacaa tgctggcctc cttgatgaat aaagatgggg ttctcatatc 1141 cgagggccaa ggcttcatga caagggagtt tctaaagagc ctgcgaaagc cttttggtga 1201 1261 ctttatggag cccaagtttg agtttgctgt gaagttcaat gcactggaat tagatgacag cgacttggca atatttattg ctgtcattat tctcagtgga gaccgcccag gtttgctgaa 1321 tgtgaagccc attgaagaca ttcaagacaa cctgctacaa gccctggagc tccagctgaa 1381 gctgaaccac cctgagtcct cacagctgtt tgccaagctg ctccagaaaa tgacagacct 1441 cagacagatt gtcacggaac acgtgcagct actgcaggtg atcaagaaga cggagacaga 1501 1561 catgagtett caccegetee tgeaggagat etacaaggae ttgtactage agagagteet 1621 gagccactgc caacatttcc cttcttccag ttgcactatt ctgagggaaa atctgaccat aagaaattta ctgtgaaaaa gcgttttaaa aagaaaaggg tttagaatat gatctatttt 1741 atgcatattg tttataaaga cacatttaca atttactttt aatattaaaa attaccatat 1801 tatgaaattg c

FIG.9B

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1	MEQPQEEAPEVREEEEKEEVAEAEGAPELNGGPQHALPSSSYTDLSRSSSPPSLLDQLQM
61	GCPGASCGSLNMECRVCGDKASGFHYGVHACEGCKGFFRRTIRMKLEYEKCERSCKIQKK
121	NRNKCQYCRFQKCLALGMSHNAIRFGRMPEAEKRKLVAGLTANEGSQYNPQVADLKAFSK
181	HIYNAYLKNFNMTKKKARSILTGKASHTAPFVIHDIETLWQAEKGLVWKQLVNGLPPYKE
241	ISVHVFYRCQCTTVETVRELTEFAKSIPSFSSLFLNDQVTLLKYGVHEAIFAMLASIVNK
301	DGLLVANGSGFVTREFLRSLRKPFSDIIEPKFEFAVKFNALELDDSDLALFIAAIILCGD
361	RPGLMNVPRVEAIQDTILRALEFHLQANHPDAQYLFPKLLQKMADLRQLVTEHAQMMQRI
421	KKTETETSLHPLLQEIYKDMY

FIG.10A

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1	gaattctgcg	gagcctgcgg	gacggcggcg	ggttggcccg	taggcagccg	ggacagtgtt
61	gtacagtgtt	ttgggcatgc	acgtgatact	cacacagtgg	cttctgctca	ccaacagatg
121	aagacagatg	caccaacgag	ggtctggaat	ggtctggagt	ggtctggaaa	gcagggtcag
181	atacccctgg	aaaactgaag	cccgtggagc	aatgatctct	acaggactgc	ttcaaggctg
241	atgggaacca	ccctgtagag	gtccatctgc	gttcagaccc	agacgatgcc	agagctatga
301	ctgggcctgc	aggtgtggcg	ccgaggggag	atcagccatg	gagcagccac	aggaggaagc
361	ccctgaggtc	cgggaagagg	aggagaaaga	ggaagtggca	gaggcagaag	gagccccaga
421	gctcaatggg	ggaccacagc	atgcacttcc	ttccagcagc	tacacagacc	tctcccggag
481	ctcctcgcca	ccctcactgc	tggaccaact	gcagatgggc	tgtgacgggg	cctcatgcgg
541	cagcctcaac	atggagtgcc	gggtgtgcgg	ggacaaggca	tcgggcttcc	actacggtgt
601	tcatgcatgt	gaggggtgca	agggcttctt	ccgtcgtacg	atccgcatga	agctggagta
661	cgagaagtgt	gagcgcagct	gcaagattca	gaagaagaac	cgcaacaagt	gccagtactg
721	ccgcttccag	aagtgcctgg	cactgggcat	gtcacacaac	gctatccgtt	ttggtcggat
781	gccggaggct	gagaagagga	agctggtggc	agggctgact	gcaaacgagg	ggagccagta
841	caacccacag	gtggccgacc	tgaaggcctt	ctccaagcac	atctacaatg	cctacctgaa
901	aaacttcaac	atgaccaaaa	agaaggcccg	cagcatcctc	accggcaaag	ccagccacac
961	ggcgcccttt	gtgatccacg	acatcgagac	attgtggcag	gcagagaagg	ggctggtgtg
1021	gaagcagttg	gtgaatggcc	tgcctcccta	caaggagatc	agcgtgcacg	tcttctaccg
1081	ctaccaatge	accacagtgg	agaccgtgcg	ggagctcact	gagttcgcca	agagcatccc
1141	cagcttcagc	agcctcttcc	tcaacgacca	ggttaccctt	ctcaagtatg	gcgtgcacga
1201	ggccatcttc	gccatgctgg	cctctatcgt	caacaaggac	gggctgctgg	tagccaacgg
1261	cagtggcttt	gtcacccgtg	agticctgcg	cagcctccgc	aaacccttca	gtgatatcat
1321	tgagcctaag	tttgaatttg	ctgtcaagtt	caacgccctg	gaacttgatg	acagtgacct
1381	ggccctattc	attgcggcca	tcattctgtg	tggagaccgg	ccaggcctca	tgaacgttcc
1441	acgggtggag	gctatccagg	acaccatcct	gcgtgccctc	gaattccacc	tgcaggccaa
1501	ccaccctgat	gcccagtacc	tcttccccaa	gctgctgcag	aagatggctg	acctgcggca
1561	actggtcaco	gagcacgccc	agatgatgca	gcggatcaag	aagaccgaaa	ccpagacctc
1621	gctgcaccct	ctgctccagg	agatctacaa	ggacatgtac	taacggcggc	acccaggcct
1681	ccctgcagac	tccaatgggg	ccagcactgg	aggggcccac	ccacatgact	tttccattga
1741	ccagctctct	tcctgtcttt	gttgtctccc	tctttctcag	ttcctctttc	ttttctaatt
1801	cctgttgcto	tgtttcttcc	tttctgtagg	tttctctctt	cccttctccc	ttctcccttg
1861	ccctcccttt	ctctctccta	tccccacgtc	: tgtcctcctt	tcttattctg	tgagatgttt
1921	tgtattatti	caccagcago	: atagaacagg	, acctctgctt	ttgcacacct	tttccccagg
1981	agcagaagag	g agtgggcctg	ccctctgccc	catcattgca	cctgcaggct	taggtcctca
2041	cttctqtctc	ctgtcttcag	agcaaaagac	ttgagccatc	: caaagaaaca	ctaagctctc
2101	tgggcctgg	g ttccagggaa	ggctaagcat	t ggcctggact	, gactgcagco	ccctatagtc
2161	atggggtcc	c tgctgcaaag	g gacagtggca	a gaccccggca	gtagagccga	gatgcctccc
2221	caagactgt	c attgccccto	cgatcgtgag	g gccacccact	: gacccaatga	tcctctccag
2281	cagcacacc	t cagccccact	gacacccagi	t gtccttccat	: cttcacactg	g gtttgccagg
2341	ccaatgttg	c tgatggccc	c tccagcaca	c acacataago	c actgaaatca	a ctttacctgc
2401	aggcaccat	g cacctcccti	t ccctccctg	a ggcaggtgag	g aacccagaga	gaggggcctg

FIG.10B

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2461	caggtgagca	ggcagggctg	ggccaggtct	ccggggaggc	aggggtcctg	caggtcctgg
2521	tgggtcagcc	cagcacctcg	cccagtggga	gcttcccggg	ataaactgag	cctgttcatt
2581	ctgatgtcca	tttgtcccaa	tagctctact	gccctcccct	tcccctttac	tcagcccagc
2641	tggccaccta	gaagtctccc	tgcacagcct	ctagtgtccg	gggaccttgt	gggaccagtc
2701	ccacaccgct	ggtccctgcc	ctcccctgct	cccaggttga	ggtgcgctca	cctcagagca
2761	gggccaaagc	acagctgggc	atgccatgtc	tgagcggcgc	agagccctcc	aggcctgcag
2821	gggcaagggg	ctggctggag	tctcagagca	cagaggtagg	agaactgggg	ttcaagccca
2881	gacttcctgg	gtcctgcctg	gtcctccctc	ccaaggagcc	attctatgtg	actctgggtg
2941	gaagtgccca	gcccctgcct	gacggnnnnn	nngatcactc	tctgctggca	ggattcttcc
3001	cactccccac	ctacccagct	gatgggggtt	ggggtgcttc	tttcagccaa	ggctatgaag
3061	ggacagetge	tgggacccac	ctccccctt	ccccggccac	atgccgcgtc	cctgccccca
3121	cccagatcta	gtgctgagga	tacagctctt	ctcagtgtct	gaacaatctc	caaaattgaa
3181	atgtatattt	ttgctaggag	ccccagcttc	ctgtgttttt	aatataaata	gtgtacacag
3241	actgacgaaa	ctttaaataa	atgggaatta	aatatttaaa	aaaaaaagcg	gccgcgaatt
3301						

FIG.10C

#### SEQUENCE LISTING .

- (1) GENERAL INFORMATION:
- (i) APPLICANT: Merck & Co., Inc.
- (ii) TITLE OF INVENTION: ASSAYS FOR NUCLEAR RECEPTOR
  AGONISTS AND ANTAGONISTS USING FLUORESCENCE RESONANCE
  ENERGY TRANSFER
- (iii) NUMBER OF SEQUENCES: 11
- (iv) CORRESPONDENCE ADDRESS:
  - (A) ADDRESSEE: Merck & Co., Inc.
  - (B) STREET: P.O. Box 2000, 126 E. Lincoln Ave.
  - (C) CITY: Rahway
  - (D) STATE: NJ

• •

- (E) COUNTRY: USA
- (F) ZIP: 07065-0900
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Diskette
  - (B) COMPUTER: IBM Compatible
  - (C) OPERATING SYSTEM: Windows
  - (D) SOFTWARE: FastSEQ for Windows Version 2.0b
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER:
  - (B) FILING DATE:
  - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
  - (A) APPLICATION NUMBER:
  - (B) FILING DATE:
- (viii) ATTORNEY/AGENT INFORMATION:
  - (A) NAME: Coppola, Joseph A
  - (B) REGISTRATION NUMBER: 38,413
  - (C) REFERENCE/DOCKET NUMBER: 20017PCT
- (ix) TELECOMMUNICATION INFORMATION:
  - (A) TELEPHONE: 732-594-6734
  - (B) TELEFAX: 732-594-4720
  - (C) TELEX:
  - (2) INFORMATION FOR SEQ ID NO:1:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 405 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

#### (ii) MOLECULE TYPE: protein

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

Met Ala Glu Asn Leu Leu Asp Gly Pro Pro Asn Pro Lys Arg Ala Lys 1 Leu Ser Ser Pro Gly Phe Ser Ala Asn Asp Ser Thr Asp Phe Gly Ser 20 Leu Phe Asp Leu Glu Asn Asp Leu Pro Asp Glu Leu Ile Pro Asn Gly Gly Glu Leu Gly Leu Leu Asn Ser Gly Asn Leu Val Pro Asp Ala Ala Ser Lys His Lys Gln Leu Ser Glu Leu Leu Arg Gly Gly Ser Gly Ser 65 Ser Ile Asn Pro Gly Ile Gly Asn Val Ser Ala Ser Ser Pro Val Gln Gln Gly Leu Gly Gln Ala Gln Gly Gln Pro Asn Ser Ala Asn Met 105 Ala Ser Leu Ser Ala Met Gly Lys Ser Pro Leu Ser Gln Gly Asp Ser 115 Ser Ala Pro Ser Leu Pro Lys Gln Ala Ala Ser Thr Ser Gly Pro Thr 135 Pro Ala Ala Ser Gln Ala Leu Asn Pro Gln Ala Gln Lys Gln Val Gly 145 150 Leu Ala Thr Ser Ser Pro Ala Thr Ser Gln Thr Gly Pro Gly Ile Cys 165 170 Met Asn Ala Asn Phe Asn Gln Thr His Pro Gly Leu Leu Asn Ser Asn 180 Ser Gly His Ser Leu Ile Asn Gln Ala Ser Gln Gly Gln Ala Gln Val 200 Met Asn Gly Ser Leu Gly Ala Ala Gly Arg Gly Arg Gly Ala Gly Met 210 215 220 Pro Tyr Pro Thr Pro Ala Met Gln Gly Ala Ser Ser Ser Val Leu Ala 225 230 235 240 Glu Thr Leu Thr Gln Val Ser Pro Gln Met Thr Gly His Ala Gly Leu 245 250 Asn Thr Ala Gln Ala Gly Gly Met Ala Lys Met Gly Ile Thr Gly Asn 260 265 Thr Ser Pro Phe Gly Gln Pro Phe Ser Gln Ala Gly Gly Gln Pro Met 275 285 Gly Ala Thr Gly Val Asn Pro Gln Leu Ala Ser Lys Gln Ser Met Val 295 300 Asn Ser Leu Pro Thr Phe Pro Thr Asp Ile Lys Asn Thr Ser Val Thr 305 310 315 320 Asn Val Pro Asn Met Ser Gln Met Gln Thr Ser Val Gly Ile Val Pro 325 330 335 Thr Gln Ala Ile Ala Thr Gly Pro Thr Ala Asp Pro Glu Lys Arg Lys 340 345 Leu Ile Gln Gln Leu Val Leu Leu His Ala His Lys Cys Gln 355 360 Arg Arg Glu Gln Ala Asn Gly Glu Val Arg Ala Cys Ser Leu Pro His 370 375 380 Cys Arg Thr Met Lys Asn Val Leu Asn His Met Thr His Cys Gln Ala 385 390 395 Gly Lys Ala Cys Gln

405

(2) INFORMATION FOR SEQ ID NO:2:

#### (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1290 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

CGAGCCCCGA	CCCCGTCCG	GGCCCTCGCC	GGCCGCGCCG	CCCGTGCCCG	GGGCTGTTTT	60
CCCGAGCAGG	TGAAAATGGC	TGAGAACTTG	CTGGACGGAC	CGCCCAACCC	CAAAAGAGCC	120
AAACTCAGCT	CGCCCGGTTT	CTCGGCGAAT	GACAGCACAG	ATTTTGGATC	ATTGTTTGAC	180
TTGGAAAATG	ATCTTCCTGA	TGAGCTGATA	CCCAATGGAG	GAGAATTAGG	CCTTTTAAAC	240
AGTGGGAACC	TTGTTCCAGA	TGCTGCTTCC	AAACATAAAC	AACTGTCGGA	GCTTCTACGA	300
GGAGGCAGCG	GCTCTAGTAT	CAACCCAGGA	ATAGGAAATG	TGAGCGCCAG	CAGCCCCGTG	360
CAGCAGGGCC	TGGGTGGCCA	GGCTCAAGGG	CAGCCGAACA	GTGCTAACAT	GGCCAGCCTC	420
AGTGCCATGG	GCAAGAGCCC	TCTGAGCCAG	GGAGATTCTT	CAGCCCCCAG	CCTGCCTAAA	480
CAGGCAGCCA	GCACCTCTGG	GCCCACCCC	GCTGCCTCCC	AAGCACTGAA	TCCGCAAGCA	540
CAAAAGCAAG	TGGGGCTGGC	GACTAGCAGC	CCTGCCACGT	CACAGACTGG	ACCTGGTATC	600
TGCATGAATG	CTAACTTTAA	CCAGACCCAC	CCAGGCCTCC	TCAATAGTAA	CTCTGGCCAT	660
AGCTTAATTA	ATCAGGCTTC	ACAAGGGCAG	GCGCAAGTCA	TGAATGGATC	TCTTGGGGCT	720
GCTGGCAGAG	GAAGGGGAGC	TGGAATGCCG	TACCCTACTC	CAGCCATGCA	GGGCGCCTCG	780
AGCAGCGTGC	TGGCTGAGAC	CCTAACGCAG	GTTTCCCCGC	AAATGACTGG	TCACGCGGGA	840
CTGAACACCG	CACAGGCAGG	AGGCATGGCC	AAGATGGGAA	TAACTGGGAA	CACAAGTCCA	900
TTTGGACAGC	CCTTTAGTCA	AGCTGGAGGG	CAGCCAATGG	GAGCCACTGG	AGTGAACCCC	960
CAGTTAGCCA	GCAAACAGAG	CATGGTCAAC	AGTTTGCCCA	CCTTCCCTAC	AGATATCAAG	1020
AATACTTCAG	TCACCAACGT	GCCAAATATG	TCTCAGATGC	AAACATCAGT	GGGAATTGTA	1080
CCCACACAAG	CAATTGCAAC	AGGCCCCACT	GCAGATCCTG	AAAAACGCAA	ACTGATACAG	1140
CAGCAGCTGG	TTCTACTGCT	TCATGCTCAT	AAGTGTCAGA	GACGAGAGCA	AGCAAACGGA	1200
GAGGTTCGGG	CCTGCTCGCT	CCCGCATTGT	CGAACCATGA	AAAACGTTTT	GAATCACATG	1260
ACGCATTGTC	AGGCTGGGAA	AGCCTGCCAA				1290

#### (2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 468 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

```
Gly Ser Val Ile Thr Asp Thr Leu Ser Pro Ala Ser Ser Pro Ser Ser
65
                    70
                                                              80
Val Thr Tyr Pro Val Val Pro Gly Ser Val Asp Glu Ser Pro Ser Gly
                85
                                     90
Ala Leu Asn Ile Glu Cys Arg Ile Cys Gly Asp Lys Ala Ser Gly Tyr
            100
                                 105
His Tyr Gly Val His Ala Cys Glu Gly Cys Lys Gly Phe Phe Arg Arg
        115
Thr Ile Arg Leu Lys Leu Val Tyr Asp Lys Cys Asp Arg Ser Cys Lys
    130
                         135
                                             140
Ile Gln Lys Lys Asn Arg Asn Lys Cys Gln Tyr Cys Arg Phe His Lys
145
                    150
                                         155
Cys Leu Ser Val Gly Met Ser His Asn Ala Ile Arg Phe Gly Arg Met
                165
                                                          175
Pro Arg Ser Glu Lys Ala Lys Leu Lys Ala Glu Ile Leu Thr Cys Glu
            180
                                 185
                                                      190
His Asp Ile Glu Asp Ser Glu Thr Ala Asp Leu Lys Ser Leu Ala Lys
                             200
Arg Ile Tyr Glu Ala Tyr Leu Lys Asn Phe Asn Met Asn Lys Val Lys
    210
                         215
Ala Arg Val Ile Leu Ser Gly Lys Ala Ser Asn Asn Pro Pro Phe Val
225
                    230
                                         235
                                                              240
Ile His Asp Met Glu Thr Leu Cys Met Ala Glu Lys Thr Leu Val Ala
                245
                                     250
Lys Leu Val Ala Asn Gly Ile Gln Asn Lys Glu Val Glu Val Arg Ile
            260
                                 265
                                                      270
Phe His Cys Cys Gln Cys Thr Ser Val Glu Thr Val Thr Glu Leu Thr
        275
                             280
                                                  285
Glu Phe Ala Lys Ala Ile Pro Ala Phe Ala Asn Leu Asp Leu Asn Asp
                         295
                                              300
Gln Val Thr Leu Leu Lys Tyr Gly Val Tyr Glu Ala Ile Phe Ala Met
305
                    310
                                                              320
Leu Ser Ser Val Met Asn Lys Asp Gly Met Leu Val Ala Tyr Gly Asn
                325
                                     330
                                                          335
Gly Phe Ile Thr Arg Glu Phe Leu Lys Ser Leu Arg Lys Pro Phe Cys
            340
                                 345
Asp Ile Met Glu Pro Lys Phe Asp Phe Ala Met Lys Phe Asn Ala Leu
                             360
                                                  365
Glu Leu Asp Asp Ser Asp Ile Ser Leu Phe Val Ala Ala Ile Ile Cys
    370
                         375
                                              380
Cys Gly Asp Arg Pro Gly Leu Leu Asn Val Gly His Ile Glu Lys Met
385
                    390
                                         395
                                                              400
Gln Glu Gly Ile Val His Val Leu Arg Leu His Leu Gln Ser Asn His
                405
                                     410
Pro Asp Asp Ile Phe Leu Phe Pro Lys Leu Leu Gln Lys Met Ala Asp
            420
                                 425
                                                      430
Leu Arg Gln Leu Val Thr Glu His Ala Gln Leu Val Gln Ile Ile Lys
        435
                             440
Lys Thr Glu Ser Asp Ala Ala Leu His Pro Leu Leu Gln Glu Ile Tyr
    450
                         455
                                              460
Arg Asp Met Tyr
465
```

#### (2) INFORMATION FOR SEQ ID NO:4:

#### (i) SEQUENCE CHARACTERISTICS:

WO 99/18124 PCT/US98/21049 .

(A) LENGTH: 1854 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

GGCCCAGGCT	GAAGCTCAGG	GCCCTGTCTG	CTCTGTGGAC	TCAACAGTTT	GTGGCAAGAC	60
	CTGAGAAGCT			TGGGAAGTTC		120
TGCCAGCAGA			TGCTTCCTGC	TTCATAGATA	AGAGTAGCTT	180
GGAGCTCGGC	GGCACAACCA	GCACCATCTG	GTCGCGATGG	TGGACACGGA	AAGCCCACTC	240
TGCCCCCTCT	CCCCACTCGA	GGCCGGCGAT	CTAGAGAGCC	CGTTATCTGA	AGAGTTCCTG	300
CAAGAAATGG	GAAACATCCA	AGAGATTTCG	CAATCCATCG	GCGAGGATAG	TTCTGGAAGC	360
TTTGGCTTTA	CGGAATACCA	GTATTTAGGA	AGCTGTCCTG	GCTCAGATGG	CTCGGTCATC	420
ACGGACACGC	TTTCACCAGC	TTCGAGCCCC	TCCTCGGTGA	CTTATCCTGT	GGTCCCCGGC	480
AGCGTGGACG	AGTCTCCCAG	TGGAGCATTG	AACATCGAAT	GTAGAATCTG	CGGGGACAAG	540
GCCTCAGGCT	ATCATTACGG	AGTCCACGCG	TGTGAAGGCT	GCAAGGGCTT	CTTTCGGCGA	600
ACGATTCGAC	TCAAGCTGGT	GTATGACAAG	TGCGACCGCA	GCTGCAAGAT	CCAGAAAAAG	660
AACAGAAACA	AATGCCAGTA	TTGTCGATTT	CACAAGTGCC	TTTCTGTCGG	GATGTCACAC	720
AACGCGATTC	GTTTTGGACG	AATGCCAAGA	TCTGAGAAAG	CAAAACTGAA	AGCAGAAATT	780
CTTACCTGTG	AACATGACAT	AGAAGATTCT	GAAACTGCAG	ATCTCAAATC	TCTGGCCAAG	840
AGAATCTACG	AGGCCTACTT	GAAGAACTTC	AACATGAACA	AGGTCAAAGC	CCGGGTCATC	900
CTCTCAGGAA	AGGCCAGTAA	CAATCCACCT	TTTGTCATAC	ATGATATGGA	GACACTGTGT	960
ATGGCTGAGA	AGACGCTGGT	GGCCAAGCTG	GTGGCCAATG	GCATCCAGAA	CAAGGAGGTG	1020
GAGGTCCGCA	TCTTTCACTG	CTGCCAGTGC	ACGTCAGTGG	AGACCGTCAC	GGAGCTCACG	1080
GAATTCGCCA	AGGCCATCCC	AGCGTTCGCA	AACTTGGACC	TGAACGATCA	AGTGACATTG	1140
CTAAAATACG	GAGTTTATGA	GGCCATATTC	GCCATGCTGT	CTTCTGTGAT	GAACAAAGAC	1200
GGGATGCTGG	TAGCGTATGG	AAATGGGTTT	ATAACTCGTG	AATTCCTAAA	AAGCCTAAGG	1260
AAACCGTTCT	GTGATATCAT	GGAACCCAAG	TTTGATTTTG	CCATGAAGTT	CAATGCACTG	1320
GAACTGGATG	ACAGTGATAT	CTCCCTTTTT	GTGGCTGCTA	TCATTTGCTG	TGGAGATCGT	1380
CCTGGCCTTC	TAAACGTAGG	ACACATTGAA	AAAATGCAGG	AGGGTATTGT	ACATGTGCTC	1440
AGACTCCACC	TGCAGAGCAA	CCACCGGAC	GATATCTTTC	TCTTCCCAAA	ACTTCTTCAA	1500
AAAATGGCAG	ACCTCCGGCA	GCTGGTGACG	GAGCATGCGC	AGCTGGTGCA	GATCATCAAG	1560
AAGACGGAGT	CGGATGCTGC	GCTGCACCCG	CTACTGCAGG	AGATCTACAG	GGACATGTAC	1620
TGAGTTCCTT	CAGATCAGCC	ACACCTTTTC	CAGGAGTTCT	GAAGCTGACA	GCACTACAAA	1680
GGAGACGGGG	GAGCAGCACG	ATTTTGCACA	AATATCCACC	ACTTTAACCT	TAGAGCTTGG	1740
					CCAGTACTTC	1800
TAAGAGCATA	GAACTCAAAT	GCTGGGGGAG	GTGGCTAATC	TCAGGACTGG	GAAG	1854

- (2) INFORMATION FOR SEQ ID NO:5:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 478 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Met Thr Met Val Asp Thr Glu Ile Ala Phe Trp Pro Thr Asn Phe Gly 1 5 10 15

Ile Ser Ser Val Asp Leu Ser Val Met Glu Asp His Ser His Ser Phe Asp Ile Lys Pro Phe Thr Thr Val Asp Phe Ser Ser Ile Ser Thr Pro His Tyr Glu Asp Ile Pro Phe Thr Arg Thr Asp Pro Val Val Ala Asp Tyr Lys Tyr Asp Leu Lys Leu Gln Glu Tyr Gln Ser Ala Ile Lys Val Glu Pro Ala Ser Pro Pro Tyr Tyr Ser Glu Lys Thr Gln Leu Tyr Asn Lys Pro His Glu Glu Pro Ser Asn Ser Leu Met Ala Ile Glu Cys Arg Val Cys Gly Asp Lys Ala Ser Gly Phe His Tyr Gly Val His Ala Cys Glu Gly Cys Lys Gly Phe Phe Arg Arg Thr Ile Arg Leu Lys Leu Ile Tyr Asp Arg Cys Asp Leu Asn Cys Arg Ile His Lys Lys Ser Arg Asn Lys Cys Gln Tyr Cys Arg Phe Gln Lys Cys Leu Ala Val Gly Met Ser His Asn Ala Ile Arg Phe Gly Arg Ile Ala Gln Ala Glu Lys Glu Lys Leu Leu Ala Glu Ile Ser Ser Asp Ile Asp Gln Leu Asn Pro Glu Ser Ala Asp Leu Arg Gln Ala Leu Ala Lys His Leu Tyr Asp Ser Tyr Ile Lys Ser Phe Pro Leu Thr Lys Ala Lys Ala Arg Ala Ile Leu Thr Gly Lys Thr Thr Asp Lys Ser Pro Phe Val Ile Tyr Asp Met Asn Ser Leu Met Met Gly Glu Asp Lys Ile Lys Phe Lys His Ile Thr Pro Leu Gln Glu Gln Ser Lys Glu Val Ala Ile Arg Ile Phe Gln Gly Cys Gln Phe Arg Ser Val Glu Ala Val Gln Glu Ile Thr Glu Tyr Ala Lys Ser Ile Pro Gly Phe Val Asn Leu Asp Leu Asn Asp Gln Val Thr Leu Leu Lys Tyr Gly Val His Glu Ile Ile Tyr Thr Met Leu Ala Ser Leu Met Asn Lys Asp Gly Val Leu Ile Ser Glu Gly Gln Gly Phe Met Thr Arg Glu Phe Leu Lys Ser Leu Arg Lys Pro Phe Gly Asp Phe Met Glu Pro Lys Phe Glu Phe Ala Val Lys Phe Asn Ala Leu Glu Leu Asp Asp Ser Asp Leu Ala Ile Phe Ile Ala Val Ile Ile Leu Ser Gly Asp Arg Pro Gly Leu Leu Asn Val Lys Pro Ile Glu Asp Ile Gln Asp Asn Leu Leu Gln Ala Leu Glu Leu Gln Leu Lys Leu Asn His Pro Glu Ser Ser Gln Leu Phe Ala Lys Leu Leu Gln Lys Met Thr Asp Leu Arg Gln Ile Val Thr Glu His Val Gln Leu Leu Gln Val Ile Lys Lys Thr Glu Thr Asp Met 

Ser Leu His Pro Leu Leu Gln Glu Ile Tyr Lys Asp Leu Tyr 465 470 475

- (2) INFORMATION FOR SEQ ID NO:6:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 1811 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

CCGACCTTAC	CCCAGGCGGC	CTTGACGTTG	GTCTTGTCGG	CAGGAGACAG	CACCATGGTG	60
GGTTCTCTCT	GAGTCTGGGA	ATTCCCGAGC	CCGAGCCGCA	GCCGCCGCCT	GGGGGGCTTG	120
GGTCGGCCTC	GAGGACACCG	GAGAGGGGCG	CCACGCCGCC	GTGGCCGCAG	AAATGACCAT	180
GGTTGACACA	GAGATCGCAT	TCTGGCCCAC	CAACTTTGGG	ATCAGCTCCG	TGGATCTCTC	240
CGTAATGGAA	GACCACTCCC	ACTCCTTTGA	TATCAAGCCC	TTCACTACTG	TTGACTTCTC	300
CAGCATTTCT	ACTCCACATT	ACGAAGACAT	TCCATTCACA	AGAACAGATC	CAGTGGTTGC	360
AGATTACAAG	TATGACCTGA	AACTTCAAGA	GTACCAAAGT	GCAATCAAAG	TGGAGCCTGC	420
ATCTCCACCT	TATTATTCTG	AGAAGACTCA	GCTCTACAAT	AAGCCTCATG	AAGAGCCTTC	480
CAACTCCCTC	ATGGCAATTG	AATGTCGTGT	CTGTGGAGAT	AAAGCTTCTG	GATTTCACTA	540
TGGAGTTCAT	GCTTGTGAAG	GATGCAAGGG	TTTCTTCCGG	AGAACAATCA	GATTGAAGCT	600
TATCTATGAC	AGATGTGATC	TTAACTGTCG	GATCCACAAA	AAAAGTAGAA	ATAAATGTCA	660
GTACTGTCGG	TTTCAGAAAT	GCCTTGCAGT	GGGGATGTCT	CATAATGCCA	TCAGGTTTGG	720
GCGGATCGCA	CAGGCCGAGA	AGGAGAAGCT	GTTGGCGGAG	ATCTCCAGTG	ATATCGACCA	780
GCTGAATCCA	GAGTCCGCTG	ACCTCCGTCA	GGCCCTGGCA	AAACATTTGT	ATGACTCATA	840
CATAAAGTCC	TTCCCGCTGA	CCAAAGCAAA	GGCGAGGGCG	ATCTTGACAG	GAAAGACAAC	900
AGACAAATCA	CCATTCGTTA	TCTATGACAT	GAATTCCTTA	ATGATGGGAG	AAGATAAAAT	960
CAAGTTCAAA	CACATCACCC	CCCTGCAGGA	GCAGAGCAAA	GAGGTGGCCA	TCCGCATCTT	1020
TCAGGGCTGC	CAGTITCGCT	CCGTGGAGGC	TGTGCAGGAG	ATCACAGAGT	ATGCCAAAAG	1080
CATTCCTGGT	TTTGTAAATC	TTGACTTGAA	CGACCAAGTA	ACTCTCCTCA	AATATGGAGT	1140
CCACGAGATC	ATTTACACAA	TGCTGGCCTC	CTTGATGAAT	AAAGATGGGG	TTCTCATATC	1200
CGAGGGCCAA	GGCTTCATGA	CAAGGGAGTT	TCTAAAGAGC	CTGCGAAAGC	CTTTTGGTGA	1260
CTTTATGGAG	CCCAAGTTTG	AGTTTGCTGT	GAAGTTCAAT	GCACTGGAAT	TAGATGACAG	1320
CGACTTGGCA	ATATTTATTG	CTGTCATTAT	TCTCAGTGGA	GACCGCCCAG	GTTTGCTGAA	1380
TGTGAAGCCC	ATTGAAGACA	TTCAAGACAA	CCTGCTACAA	GCCCTGGAGC	TCCAGCTGAA	1440
GCTGAACCAC	CCTGAGTCCT	CACAGCTGTT	TGCCAAGCTG	CTCCAGAAAA	TGACAGACCT	1500
CAGACAGATT	GTCACGGAAC	ACGTGCAGCT	ACTGCAGGTG	ATCAAGAAGA	CGGAGACAGA	1560
CATGAGTCTT	CACCCGCTCC	TGCAGGAGAT	CTACAAGGAC	TTGTACTAGC	AGAGAGTCCT	1620
GAGCCACTGC	CAACATTTCC	CTTCTTCCAG	TTGCACTATT	CTGAGGGAAA	ATCTGACCAT	1680
AAGAAATTTA	CTGTGAAAAA	GCGTTTTAAA	AAGAAAAGGG	TTTAGAATAT	GATCTATTTT	1740
ATGCATATTG	TTTATAAAGA	CACATTTACA	ATTTACTTTT	AATATTAAAA	ATTACCATAT	1800
TATGAAATTG	С					1811

- (2) INFORMATION FOR SEQ ID NO:7:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 441 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Met Glu Gln Pro Gln Glu Glu Ala Pro Glu Val Arg Glu Glu Glu Glu Lys Glu Glu Val Ala Glu Ala Glu Gly Ala Pro Glu Leu Asn Gly Gly Pro Gln His Ala Leu Pro Ser Ser Ser Tyr Thr Asp Leu Ser Arg Ser 35 Ser Ser Pro Pro Ser Leu Leu Asp Gln Leu Gln Met Gly Cys Asp Gly Ala Ser Cys Gly Ser Leu Asn Met Glu Cys Arg Val Cys Gly Asp Lys 65 70 Ala Ser Gly Phe His Tyr Gly Val His Ala Cys Glu Gly Cys Lys Gly 85 Phe Phe Arg Arg Thr Ile Arg Met Lys Leu Glu Tyr Glu Lys Cys Glu Arg Ser Cys Lys Ile Gln Lys Lys Asn Arg Asn Lys Cys Gln Tyr Cys 115 120 Arg Phe Gln Lys Cys Leu Ala Leu Gly Met Ser His Asn Ala Ile Arg 130 140 Phe Gly Arg Met Pro Glu Ala Glu Lys Arg Lys Leu Val Ala Gly Leu Thr Ala Asn Glu Gly Ser Gln Tyr Asn Pro Gln Val Ala Asp Leu Lys 165 Ala Phe Ser Lys His Ile Tyr Asn Ala Tyr Leu Lys Asn Phe Asn Met 180 185 190 Thr Lys Lys Ala Arg Ser Ile Leu Thr Gly Lys Ala Ser His Thr 200 Ala Pro Phe Val Ile His Asp Ile Glu Thr Leu Trp Gln Ala Glu Lys 210 Gly Leu Val Trp Lys Gln Leu Val Asn Gly Leu Pro Pro Tyr Lys Glu 225 230 235 240 Ile Ser Val His Val Phe Tyr Arg Cys Gln Cys Thr Thr Val Glu Thr Val Arg Glu Leu Thr Glu Phe Ala Lys Ser Ile Pro Ser Phe Ser Ser 265 Leu Phe Leu Asn Asp Gln Val Thr Leu Leu Lys Tyr Gly Val His Glu 275 280 285 Ala Ile Phe Ala Met Leu Ala Ser Ile Val Asn Lys Asp Gly Leu Leu 290 295 300 Val Ala Asn Gly Ser Gly Phe Val Thr Arg Glu Phe Leu Arg Ser Leu 305 310 315 Arg Lys Pro Phe Ser Asp Ile Ile Glu Pro Lys Phe Glu Phe Ala Val 325 330 335 Lys Phe Asn Ala Leu Glu Leu Asp Asp Ser Asp Leu Ala Leu Phe Ile 340 345 350 Ala Ala Ile Ile Leu Cys Gly Asp Arg Pro Gly Leu Met Asn Val Pro 360 Arg Val Glu Ala Ile Gln Asp Thr Ile Leu Arg Ala Leu Glu Phe His 370 375 380 Leu Gln Ala Asn His Pro Asp Ala Gln Tyr Leu Phe Pro Lys Leu Leu 385 390 395 Gln Lys Met Ala Asp Leu Arg Gln Leu Val Thr Glu His Ala Gln Met 405 410 415

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Met Gln Arg Ile Lys Lys Thr Glu Thr Glu Thr Ser Leu His Pro Leu 420 425 430

Leu Gln Glu Ile Tyr Lys Asp Met Tyr 435 440

#### (2) INFORMATION FOR SEQ ID NO:8:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 3301 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

GAATTCTGCG	GAGCCTGCGG	GACGGCGCG	GGTTGGCCCG	TAGGCAGCCG	GGACAGTGTT	60
GTACAGTGTT	TTGGGCATGC	ACGTGATACT	CACACAGTGG	CTTCTGCTCA	CCAACAGATG	120
AAGACAGATG	CACCAACGAG	GGTCTGGAAT	GGTCTGGAGT	GGTCTGGAAA	GCAGGGTCAG	180
ATACCCCTGG	AAAACTGAAG	CCCGTGGAGC	AATGATCTCT	ACAGGACTGC	TTCAAGGCTG	240
ATGGGAACCA	CCCTGTAGAG	GTCCATCTGC	GTTCAGACCC	AGACGATGCC	AGAGCTATGA	300
CTGGGCCTGC	AGGTGTGGCG	CCGAGGGGAG	ATCAGCCATG	GAGCAGCCAC	AGGAGGAAGC	360
CCCTGAGGTC	CGGGAAGAGG	AGGAGAAAGA	GGAAGTGGCA	GAGGCAGAAG	GAGCCCCAGA	420
GCTCAATGGG	GGACCACAGC	ATGCACTTCC	TTCCAGCAGC	TACACAGACC	TCTCCCGGAG	480
CTCCTCGCCA	CCCTCACTGC	TGGACCAACT	GCAGATGGGC	TGTGACGGGG	CCTCATGCGG	540
CAGCCTCAAC	ATGGAGTGCC	GGGTGTGCGG	GGACAAGGCA	TCGGGCTTCC	ACTACGGTGT	600
TCATGCATGT	GAGGGGTGCA	AGGGCTTCTT	CCGTCGTACG	ATCCGCATGA	AGCTGGAGTA	660
CGAGAAGTGT	GAGCGCAGCT	GCAAGATTCA	GAAGAAGAAC	CGCAACAAGT	GCCAGTACTG	720
CCGCTTCCAG	AAGTGCCTGG	CACTGGGCAT	GTCACACAAC	GCTATCCGTT	TTGGTCGGAT	780
GCCGGAGGCT	GAGAAGAGGA	AGCTGGTGGC	AGGGCTGACT	GCAAACGAGG	GGAGCCAGTA	840
CAACCCACAG	GTGGCCGACC	TGAAGGCCTT	CTCCAAGCAC	ATCTACAATG	CCTACCTGAA	900
AAACTTCAAC	ATGACCAAAA	AGAAGGCCCG	CAGCATCCTC	ACCGGCAAAG	CCAGCCACAC	960
GGCGCCCTTT	GTGATCCACG	ACATCGAGAC	ATTGTGGCAG	GCAGAGAAGG	GGCTGGTGTG	1020
GAAGCAGTTG	GTGAATGGCC	TGCCTCCCTA	CAAGGAGATC	AGCGTGCACG	TCTTCTACCG	1080
CTGCCAGTGC	ACCACAGTGG	AGACCGTGCG	GGAGCTCACT	GAGTTCGCCA	AGAGCATCCC	1140
CAGCTTCAGC	AGCCTCTTCC	TCAACGACCA	GGTTACCCTT	CTCAAGTATG	GCGTGCACGA	1200
GGCCATCTTC	GCCATGCTGG	CCTCTATCGT	CAACAAGGAC	GGGCTGCTGG	TAGCCAACGG	1260
CAGTGGCTTT	GTCACCCGTG	AGTTCCTGCG	CAGCCTCCGC	AAACCCTTCA	GTGATATCAT	1320
TGAGCCTAAG	TTTGAATTTG	CTGTCAAGTT	CAACGCCCTG	GAACTTGATG	ACAGTGACCT	1380
GGCCCTATTC	ATTGCGGCCA	TCATTCTGTG	TGGAGACCGG	CCAGGCCTCA	TGAACGTTCC	1440
ACGGGTGGAG	GCTATCCAGG	ACACCATCCT	GCGTGCCCTC	GAATTCCACC	TGCAGGCCAA	1500
CCACCCTGAT	GCCCAGTACC	TCTTCCCCAA	GCTGCTGCAG	AAGATGGCTG	ACCTGCGGCA	1560
ACTGGTCACC	GAGCACGCCC	AGATGATGCA	GCGGATCAAG	AAGACCGAAA	CCGAGACCTC	1620
					ACCCAGGCCT	1680
CCCTGCAGAC	TCCAATGGGG	CCAGCACTGG	AGGGGCCCAC	CCACATGACT	TTTCCATTGA	1740
					TTTTCTAATT	1800
CCTGTTGCTC	TGTTTCTTCC	TTTCTGTAGG	TTTCTCTCTT	CCCTTCTCCC	TTCTCCCTTG	1860
CCCTCCCTTT	CTCTCTCTA	TCCCCACGTC	TGTCCTCCTT	TCTTATTCTG	TGAGATGTTT	1920
TGTATTATTT	CACCAGCAGC	ATAGAACAGG	ACCTCTGCTT	TTGCACACCT	TTTCCCCAGG	1980
	• <b>-</b>			<del>-</del>	TAGGTCCTCA	2040
CTTCTGTCTC	CTGTCTTCAG	AGCAAAAGAC	TTGAGCCATC	CAAAGAAACA	CTAAGCTCTC	2100
		GGCTAAGCAT				2160
					GATGCCTCCC	2220
					TCCTCTCCAG	2280
CAGCACACCT	CAGCCCCACT	GACACCCAGT	GTCCTTCCAT	CTTCACACTG	GTTTGCCAGG	2340

CCAATGTTGC	TGATGGCCCC	TCCAGCACAC	ACACATAAGC	ACTGAAATCA	CTTTACCTGC	2400
AGGCACCATG	CACCTCCCTT	CCCTCCCTGA	GGCAGGTGAG	AACCCAGAGA	GAGGGGCCTG	2460
CAGGTGAGCA	GGCAGGGCTG	GGCCAGGTCT	CCGGGGAGGC	AGGGGTCCTG	CAGGTCCTGG	2520
TGGGTCAGCC	CAGCACCTCG	CCCAGTGGGA	GCTTCCCGGG	ATAAACTGAG	CCTGTTCATT	2580
CTGATGTCCA	TTTGTCCCAA	TAGCTCTACT	GCCCTCCCCT	TCCCCTTTAC	TCAGCCCAGC	2640
TGGCCACCTA	GAAGTCTCCC	TGCACAGCCT	CTAGTGTCCG	GGGACCTTGT	GGGACCAGTC	2700
CCACACCGCT	GGTCCCTGCC	CTCCCCTGCT	CCCAGGTTGA	GGTGCGCTCA	CCTCAGAGCA	2760
GGGCCAAAGC	ACAGCTGGGC	ATGCCATGTC	TGAGCGGCGC	AGAGCCCTCC	AGGCCTGCAG	2820
GGGCAAGGGG	CTGGCTGGAG	TCTCAGAGCA	CAGAGGTAGG	AGAACTGGGG	TTCAAGCCCA	2880
GGCTTCCTGG	GTCCTGCCTG	GTCCTCCCTC	CCAAGGAGCC	ATTCTATGTG	ACTCTGGGTG	2940
GAAGTGCCCA	GCCCCTGCCT	GACGGNNNNN	NNGATCACTC	TCTGCTGGCA	GGATTCTTCC	3000
CGCTCCCCAC	CTACCCAGCT	GATGGGGGTT	GGGGTGCTTC	TTTCAGCCAA	GGCTATGAAG	3060
GGACAGCTGC	TGGGACCCAC	CTCCCCCCTT	CCCCGGCCAC	ATGCCGCGTC	CCTGCCCCCA	3120
CCCGGGTCTG	GTGCTGAGGA	TACAGCTCTT	CTCAGTGTCT	GAACAATCTC	CAAAATTGAA	3180
ATGTATATTT	TTGCTAGGAG	CCCCAGCTTC	CTGTGTTTTT	AATATAAATA	GTGTACACAG	3240
ACTGACGAAA	CTTTAAATAA	ATGGGAATTA	AATATTTAAA	AAAAAAAGCG	GCCGCGAATT	3300
С						3301
	^ ·		TO 110 0			

- (2) INFORMATION FOR SEQ ID NO:9:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 38 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

#### ACTCGGATCC AAGCCATGGC TGAGAACTTG CTGGACGG

38

- (2) INFORMATION FOR SEQ ID NO:10:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 32 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

#### CACAAAGCTT AGGCCATGTT AGCACTGTTC GG

32

- (2) INFORMATION FOR SEQ ID NO:11:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 37 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

CTCAGTCGAC TTATTGAATT CCACTAGCTG GAGATCC

37

# INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/21049

A. CLASSIFICATION OF SUBJECT MATTER  IPC(6) :C07K 14/435, 14/705; C09K 11/06; G01N 33/ US CL :252/301.16, 301.36, 301.4R; 435/7.8; 530/350  According to International Patent Classification (IPC) or to	. 358	
B. FIELDS SEARCHED		
Minimum documentation searched (classification system for	ollowed by classification symbols)	
U.S.: 252/301.16, 301.32, 301.4R; 435/7.8; 530/350	), 358	
	to the design of the landed	in the fields searched
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Electronic data base consulted during the international sea	arch (name of data base and, where practicable,	search terms used)
C. DOCUMENTS CONSIDERED TO BE RELEVA	ANT	
Category* Citation of document, with indication, w	here appropriate, of the relevant passages	Relevant to claim No.
coactivators: characterization b	ptors have distinct affinities for y fluorescence resonance energy ogy. October 1998. Vol. 12, No. page 1596 and figures 1-4.	1-27
Further documents are listed in the continuation of	of Box C. See patent family annex.	
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## INTERNATIONAL SEARCH REPORT

International application No. . PCT/US98/21049

	B. FIELDS SEARCHED  Electronic data bases consulted (Name of data base and where practicable terms used):					
	APS, STN, BIOSCIENCE, BIOSIS, CAPLUS, SCISEARCH					
	search terms: nuclear receptor, steroid receptor, retinoic acid receptor, co-activator, fret					
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